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A NOVEL CATALYTIC FORMULATION AND ITS PREPARATION

TIELD OF THE INVENTION

The present invention relates to a new class of heterogeneous catalysts, the methodology for which is useful in preparing solid catalysts for a variety of chemical reactions. Particularly, this invention relates to a catalyst system comprising of chemical formulation constituting an insoluble material having desired catalytic properties and support, assembled together by a specific technique. The said catalyst is useful for promoting reactions in gas or liquid phase. The unique feature of this catalyst system is that entire catalytic formulation remains as a composite solid material without disassembling during the course of reaction. The invention primarily describes a technique whereby soluble catalyst is converted to insoluble material by appropriate molecular modification. The invention further relates to preferred methods for preparation of such catalytic formulations.

BACKGROUND OF THE INVENTION

Soluble molecular catalysts, particularly complexes of transition metals are well known in the art. Such catalysts are also known to catalyze a variety of useful organic transformations. These transformations for instance include hydrogenation, hydroformylation, carbonylation, amination, isomerization, telomerization, Heck olefination, Suzuki coupling, metathesis, epoxidation etc. Such transformations find a variety of useful applications for the synthesis of pharmaceuticals, pesticides, solvents and other valuable products of industrial and consumer significance.

Amongst the established practices known in the prior art, catalytically active transition metal complexes have principally been applied in homogeneous form, as solution in a reactant phase. For example, in case of hydroformylation of olefins using rhodium and phosphine ligand complex catalyst wherein phosphine ligand is free of ionic charge such as tributyl phosphine, triphenyl phosphine etc. and soluble in the reaction medium. Although such catalysts are highly effective, in terms of productivity and selectivity, its applicability on practical grounds is often limited to volatile products. In case of reactions catalyzed by homogeneous catalysts involving high molecular weight and especially nonvolatile products catalyst separation is a critical problem. High cost of catalyst, susceptibility to high temperatures and stringent product specification demand quantitative catalyst separation. Common unit operations such as distillation and

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crystallization are least significant since, organometallic complexes being delicate in nature and cannot withstand separation stresses especially thermal stresses as encountered in distillation. Other separation techniques being inefficient in separating such a small quantity of catalyst cannot be used in effective manner. Moreover high purity of the product is of importance in products such as pharmaceuticals, demanding rigorous separation of catalyst from product stream. Thus use of homogeneous catalyst as such has suffered from inherent difficulties in the recovery of the catalysts from reaction products. Efficient catalyst recovery and recycle is the pivotal issue for the economic viability of the process since, complexes and ligands are often expensive.

It is also known in the art to use aqueous solutions of sulfonated aryl phosphines and many other water-soluble compounds and transition metal complex catalyst derived from it to effect reactions. As disclosed in patent (U.S. patent No. 4,248,802) all such reactions are operated in biphasic conditions wherein catalyst phase is aqueous and products and reactants dissolved in organic phase. Similarly reverse biphasic techniques are also applicable wherein catalyst is dissolved in organic phase and product and reactants in aqueous phase. A judicious choice is necessary while utilizing biphasic catalytic systems depending upon solubility of reactants and products. In either case at the end of reaction catalyst and product phases are separated wherefore catalyst phase is recycled and product phase is directed for further downstream processing.

It is however recognized that catalytic activity is low in biphasic medium due to limited solubility of organic reactants in the catalyst phase. Moreover such biphasic reactions require high reactor pressure in case of gas-liquid reactions. To achieve practical rates of reactions catalyst loading has to be increased or alternatively using larger process equipment, which is usually cost prohibitive. Further, these reactions require numerous accessory equipment's to separate liquid-liquid fractions under reaction conditions.

Over the past quarter of century many attempts have been made to heterogenize this versatile class of soluble catalysts. Several methods were developed with central theme being retention of high activity and selectivity as that of native catalytic species and facilitate separation by simple filtration, centrifugation or gravity settling.

One of the techniques to form a solid catalyst involves interaction of metal salt or precursor complex with solid support that is appropriately functionalized with organic functional groups that are capable of forming coordination bonds with metal. The support used in this context is either organic-polymeric or inorganic matrix. These supports are

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chemically functionalized to bear amino, phosphino and carboxylato functional groups on the surface of the support. Work related to this technique is described in, *Catalysis Reviews*, 16, 17-37 (1974); *Chemical Reviews*, 81, 109, (1981); *Tetrahedron Asymmetry*, 6, 1109-1116 (1995); *Tetrahedron Letters*, 37, 3375-3378 (1996). "Catalysis by supported complexes", Studies in surface science and catalysis, volume 8, Elsevier Publishing Co. Amsterdam, 1981 describes the complexes grafted to inorganic supports.

From practical stand point these catalysts are not widely used since their activities are frequently lower than corresponding homogeneous catalysts in addition there are various complications that are inherent due to polymeric nature of the support for example swelling and shrinking of the matrix, which alters diffusion resistance. It is also found that in long run and upon exposure to oxygen metal attached to support is lost in the solution thereby degeneration of the activity of the catalyst.

Supported liquid phase catalyst such as those described in US patent 3,855,307 (1974) and US patent 4,994,427 (1991) are critically sensitive to the character of the reaction medium and are often leached in to reaction medium depending upon the nature of the solvent. The applicability of such catalyst is limited to only vapor phase reactions. The technique as described in US patent 4,994,427(1991) wherein solution of water-soluble catalyst is distributed on high surface area solid. The aqueous film of catalyst containing solution remains insoluble in nonpolar organic phase thus, after reaction solid catalyst can be recovered by simple filtration. Applicability of such catalyst is limited to reactions involving water insoluble reaction media. Moreover such catalysts are sensitive to content of water.

Entrapment of the catalyst in porous material such as zeolite has been described by Balkus, et al in J. Inclusion Phenom. Mol. Recognit. Chem., 21(1-4), 159-84 (English) 1995 The catalyst is encapsulated in three-dimensional network of zeolites wherein, catalyst because of size exclusion can not diffuse out of zeolite but smaller sized reactants diffuse inside the zeolite and products formed subsequently diffuse out. Yet another article J. Catal, 163(2), 457-464 1996 have described the method to entrap catalyst within the polymer matrix but because of diffusion resistance, catalyst efficiency is doubtful.

Despite several known techniques for heterogenization of soluble molecular catalysts there is no known method, which can be conveniently used for diversity of catalytic entities using a common protocol. Furthermore catalyst formed by such protocol is required to provide a solid catalyst that can be used for polar as well as nonpolar

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reaction media. Certainly a particular need exists for such technique of catalyst formulation and present invention is aimed to fulfill these needs.

DISCLOSURE OF THE INVENTION

Importantly in the general as well as specific background of the art there is no teaching or suggestion of heterogeneous catalyst analogous to supported metal catalyst wherein, catalytically active material is physically distributed on the solid surface and the formulation as a whole can be employed as heterogeneous catalyst which is useful for catalyzing reactions in polar as well as nonpolar solvents. Thus it is an object of this invention to provide a novel catalyst useful for promoting a variety of chemical reactions. More particularly, this invention relates to a catalyst system comprising of calcium, strontium, barium salt of ligand containing at least two or more acidic functional groups and an organometallic catalyst generated from it. These salts are supported on the solid surface of inert vehicle or carrier. This catalyst is useful for promoting reactions in aqueous, polar and non-polar organic mediums.

Many anionically charged phosphines, and other coordinating compounds as well as variety of their salts are known in the art. It is also known that these ligands and complexes thereof are water-soluble but importantly there is no disclosure or suggestion in open literature, patent or any known reference, indicating an appreciation of any significance, of the formation of insoluble material as alkaline earth metal salts of anionically charged ligands and complexes thereof. Moreover there is no teaching, disclosure or suggestion in any reference known to applicants, evidencing any significance of particular type, class or characteristic of such insoluble organometallic complexes or catalytic application thereof as it relates directly to recovery and recycle of the catalyst.

It has now been discovered that reactions that are catalyzed otherwise by soluble catalyst can be catalyzed by solid catalyst of this invention. The solid catalyst as said herein is not a chemically defined single component catalyst system but a formulation wherein solid support and catalytically active material are assembled together to form a solid catalyst. The support is components, which by itself is a catalytically inactive but provides a physical vehicle, filler and provide a high surface area whereupon catalytically active material is placed. This conglomerate of support and catalytically active material is not a simple random physical mixture but assembled in a specific manner such that

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support surface is covered or deposited with catalytically active material. Such concepts are known earlier as described in the background of the invention but providing a catalyst suitable only for gas phase or for specific liquid phases. For example supported aqueous phase catalysts (hereafter termed as SAPC) or supported liquid phase catalysts (hereafter termed as SLPC). SAPC for instance can only be employed in cases where reaction medium is water immiscible organic media. Similarly SLPC are suitable only in gas phase but not in general liquid phases.

The surprising element of this invention is that a generic technique is discovered whereby native catalyst, which is otherwise soluble, can be converted to a solid material, which is practically insoluble in organic and aqueous medium. Catalytically active material as said herein is constructed from secondary construction blocks that are derived from catalytically active moieties, when placed over a high surface area solid, catalyze a reaction which is otherwise catalyzed by native building species in homogeneous phase but at the same time remaining as solid placed on the support. Due to such reason catalyst as a whole can be separated from reaction mixture by simple solid-liquid separation.

Such catalytic formulation provides a tremendous advantage than catalyzing a reaction by homogeneous or heterogeneous catalyst. This catalyst was conceived in manner analogous to supported heterogeneous catalyst but supported active phase is constructed from molecular entities, which in reality catalyze actual reaction. This particular formulation synergistically combines the desired facile separation and high specificity of the molecular catalysts. The advantages that were obvious to the inventors are;

- (a) Solid catalyst providing inherent separation
- (b) Activity and selectivity similar to soluble molecular catalysts since active sites are structurally isotropic
- (c) Formulation as a whole is mechanically robust material
- (d) Modularity of the assembly is such that desired selecting entities, support and additives depending on the need one can assemble the catalyst.

As described herein central theme of this discovery is the invention whereby a soluble catalytic material is converted to a solid that is practically insoluble in diversity of liquid media by a systematic molecular modification. It was realized that when soluble catalytic species is modified in such a way that it bears two or more anionic groups existing along with proton, alkali metal ammonium and quaternary ammonium salts. The soluble catalyst modification mentioned here implies that anionic functional groups are

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introduced while synthesizing components of catalyst or otherwise modifying catalyst as such. Said anionically functionalized salts when interacted with group IIA metal salts provide a solid material that is insoluble in variety of liquid media. This solid material is composed of building blocks of catalytic entities bridged with group IIA metal cation.

Surprisingly in previous patents (US 4,248,802 and US 4,994,427) alkali earth metal salts of such anionically functionalized compounds were claimed in general as aqueous soluble. In this invention disclosure we disclose that alkaline earth metal salts of said anionically functionalized compounds are insoluble in organic media or sparingly soluble or insoluble in aqueous media. Therefore in order to suppress aqueous solubility admixture of catalytically inactive insoluble salts is employed. This admixture is primarily intended to suppress solubility of ionic solids by phenomenon commonly known as common ion effect.

In a manner described earlier wide diversity of catalytic complexes can be converted in to solid material by a common protocol. Such solid materials are found to be stable under commonly encountered reaction environments. In another respect soluble catalysts for diverse classes of reactions, such as for instance hydrogenation, hydroformylation, carbonylation, olefination, telomerization, isomarization oxidation etc. can be solidified. Yet another aspect of the present invention is the formulation of this material and a solid support to form a catalyst. The support involved here can be chosen independent of catalytic entity being formulated and catalytically inactive additive that is admixed. The most interesting aspect of the present invention is that said catalytic formulation alternatively termed as catalytic ensemble or catalytic assembly, remain as a solid without its components being disintegrated by dissolution. Said ensemble can be employed for catalyzing chemical reactions in slurry or fixed bed reactor configurations.

Thus the precise object of the present invention is to provide a solid catalyst wherein catalytic entities responsible are molecularly defined and isotropic species. More over technique of synthesis should be common set of techniques whereby desired catalytic species can be heterogenized by simple means. The essential object of the present invention is that catalytically active solid formulation should not disintegrate or disassemble under the conditions of reaction as well as under liquid flow. Another desired but not essential object is to provide a solid catalyst that chemically imitates the its soluble analogue but at the same time providing facile separation due to inherent advantages of solid catalysts. The term 'native' used to mean in this context as a catalytic entity before modification and interaction with group IIA metal salts to yield a solid.

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Other objects and advantages of this invention will become readily apparent from the following written description and appended claims.

Accordingly, generic aspect of the invention can be described as the discovery of a common technique whereby a solid catalyst can be prepared. The family of catalysts that are similar in composition, feature and advantages are referred here by the term generic. The feature of this family of catalyst is that these are heterogeneous but active sites are chemically defined organometallic entities physically existing as solid. These organometallic entities are analogues derived from equivalent homogeneous catalytic entities. Homogeneous catalytic entities referred herein encompass entire class of soluble catalysts. These native structures are chemically modified to introduce negatively charged functional groups such as -SO₃⁻, -PO₃²⁻ or -COO⁻. When such material are synthesized, they exists as soluble salts depending on the counter ion accompanying anionic functional group. The most intriguing phenomenon realized in this invention itself qualifies to term invention as generic, which is organometallic entities modified as described earlier can be converted to a solid material by interacting them with group IIA metal salts. The solid formed is ultimately a salt of group IIA metals this observation is validated by converting large diversity of chemical structures to insoluble solids as described earlier moreover methods have been developed to assemble such solids on the surface of the supports.

Brief description of the drawings

Figure 1 is the schematic representation of the conceptual representation of the catalyst formulation

Figure 2 is the semi realistic enlarged view of the catalyst formulation surface. A support upon which catalytically active material is deposited, multiple or single reactants arrive to this catalytic material, which contains active site whereupon reactants are transformed in to products and released back in to bulk liquid

Figure 3 is the schematic of the continuos liquid extractor for solids wherein, a is the unidirectional gas bubbler connected to condenser, b is the condenser, c is the extraction vessel holding magnetic needle and solid to be leached/extracted, d is the magnetic stirrer unit, e is the vessel holding extraction liquid, f is the high temperature bath

Figure 4 is the schematic of the fluidized bed in which catalyst formulation is processed wherein, a is the jacket through which constant temperature fluid is

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circulated, **b** is the atomizer through which liquids are sprayed in the fluidized bed, **c** is the gas solid separation mesh, **d** is the inlet for solution A, **e** is the inlet for solution B, V1 and V2 are valves

Figure 5 is the schematic of the catalyst preparation unit with simultaneous removal of liquid wherein, a is the inert gas inlet, b is the inlet for solutions A and B, c is the vessel holding magnetic needle, support and liquid, d inert gas outlet, e is the condenser, f is the liquid collector, g is the collection arm for liquid.

Figure 6 is the schematic of the catalyst preparation unit wherein, a is the inlet for solution A and B, b is the vacuum line, c motor for coating pan, d is the coating pan, e is the nozzle for liquids A and B, f is the high temperature bath, g is the collection vessel for condensed liquid

DESCRIPTION OF THE PREFERRED EMBODIMENTS

For the purposes of promoting an understanding of the principles of the invention, reference will now be made to several embodiments illustrated in the examples and specific description will be made to describe the same. It will nevertheless be understood that no limitation of the scope of the invention is thereby intended, such alterations and further modifications in these embodiments, and such further applications of the principles of the invention as described herein being contemplated as would normally occur to one skilled in the art to which this invention relates.

In one embodiment of the inventiuon, provides a novel heterogeneous catalytic composition comprising a solid support having deposited thereon a catalytically active material which is practically insoluble in variety of liquid media, the said solid material consisting of catalytically active anionic entities with group IIA metal ions and the catalytic active material is molecularly well defined.

In another embodiment of the invention, the catalytically active entity is deposited on the external and the pore surfaces of the solid support, pores of which are predominantly of diameter greater than about $20~\text{A}^0$ and the pores of solid support having a pore diameters ranging from about $3-3000~\text{A}^0$.

In still another embodiment, the solid support is chemically inactive solid material, exists as powder, granules, flakes or pallets of regular or irregular shapes, sheets,

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monolith, ropes and woven fabric of fibrous solids and the porous solid support is mechanically robust and thermally stable solid, insoluble in reaction media.

In still another embodiment, the catalytically active entity is insoluble in reaction media, which are selected from organic, aqueous, <u>flours</u>, non-aqueous ionic liquids and supercritical fluid phases and is thermally stable solid material having melting point greater than 100 °C.

In yet another embodiment, the catalytically active material is a non-subliming solid.

In yet another embodiment of the invention providesd a catalyst comprising of solid support having deposited thereon catalytically active entity which remains as a stable composite solid in gas, liquid and gas-liquid phases and the liquid phase is selected from organic, aqueous, flours, non-aqueous ionic liquids and supercritical fluid phases or mixture thereof containing reactants, products and promoters.

In yet another embodiment, the catalyst remains as a physically stable composite solid in gas or liquid phases over a temperature range of -78 to 300 °C and over pressure ranging from 5 to 5000 psi.

In yet another embodiment, group IIA metal used is a cation having +2 charge and is selected from calcium, strontium, barium and mixtures thereof.

In yet another embodiment, group IIA metal used is selected independently or in combination with other group IIA metals.

In yet another embodiment, thecatalytically active entity is an anion having two or more negative charges and is independently selected from metal complexes, quaternary compounds, metaloxoanions, polyoxometallates and combinations thereof.

In yet another embodiment provides a catalyst wherein, the metal complexes having a general formula

$(M)_x(L)_v(L^*)_z$

wherein M is catalytic metal atom or ion of a coordination complex is a transition metal from group IIIB, IVB, VB, VIB, VIB, IB or IIB of the periodic table of elements

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and is selected independently, x is ranging from 1 to 60, L is selected from aliphatic, aromatic and heterocyclic compound containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃,-SO₂ -PO₃², -COO, -O, AsO₃² and -S⁻, y is at least 1, L* is a radical selected from organic anion, inorganic anion and coordinating compound containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene, =C: having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl and Z is ranging from 0 to 7.

In yet another embodiment, the quaternary compound is having a general formula

$[(Y^{+})(R^{*})_{I}][Z^{-}]$

wherein, I = 4 for $Y^+ = N^+$, P^+ , As^+ ; I = 3 for $Y^+ = S^+$ and R^* is selected independently from alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from $-SO_3^-$, $-SO_2^-$, $-COO^-$, $-O^-$, AsO_3^{2-} and $-S^-$ and Z is an anion selected from organic anion, inorganic anion and coordination complex anion.

In yet another embodiment, the insoluble catalytically active material optionally comprising catalytically inert additive, inert additive is an anion having two or more negative charges and which is independently selected from organic, inorganic anions or in combination thereof.

In yet another embodiment, the catalytically inert additive is selected from ligand compounds wherein, ligand compounds contain at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃-,-SO₂--PO₃²-, -COO-, -O-, AsO₃²- and -S².

In yet another embodiment, the amount of catalytically active entity employed is 40 % weight or less and the amount of catalytically inert additive employed is in the proportion of 0 to 200-weight % of catalytically active entity.

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In yet another embodiment, the catalyst can be employed to catalyze reactions in gas phase or in slurry phase.

In yet another embodiment, the catalyst further comprising a film of high boiling liquid deposited on the solid catalyst.

As stated herein above, primary aspect of applicants' invention is directed to a solid formulation as a catalyst comprising of a solid support having deposited thereon a catalytically active solid material. Thus the catalytic formulation referred herein is primarily a solid in physical sense and an organized ensemble of chemical components put together to perform the task of catalyzing the reaction. The definition of assembly is narrowly put down in embodiments that will follow, to conceive a real life catalyst. This multi-component catalyst is the balanced compromise between supposed fluid flow around catalytic particle, activity and a physical integrity. The relative importance of these factors directly affects the reaction, reactor design, process conditions and economics. Although many catalytic materials are composed of single components such as zeolites, pillared clays, metals alloys and metal oxides they certainly cannot catalyze the wide diversity of reactions. Whereas multi-component catalysts as considered in this embodiment offer a choice of physicochemical properties that can be selected from different materials as salts, oxides, metal aggregates or organometallic materials. In order to achieve earlier stated features, this embodiment describes the general architectural draft of catalytic formulation of this invention as illustrated in the figure 1 and 2

Referring to the figure detailed in figure 1 components of the said formulation is distinguished into two categories. These two categories are catalytically passive and catalytically active. Passive components are supports and other additives that necessarily impart solid character to catalyst formulation and may be selected independent of active entity depending upon application. Of course it is understood that choice of supports cannot be made randomly and selection is totally dependent on the process. For example silica supports cannot be utilized in strong alkali solutions, as it will dissolve causing the loss of integrity. Thus supports in the context of this embodiment are considered as agents to impart physical shape and form to catalyst particle and act as a vehicle to enhance the maneuverability.

From previous discussion it is explicit enough to judge that catalytically active material is often expensive and sometimes precious, later is often true for reaction that are catalyzed by organometallic complex catalysts. The activity of such materials when

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supported on solid depends on factors such as surface area, porosity, geometry and resistance to surface fouling. In an effort to optimize these factors, it is common practice to disperse active ingredients on the surface of the inactive solids conventionally known as supports or carriers. Although these materials are considered as diluents, they sometimes play an important multifunctional role in directing catalytic activity. This may include chemical reaction with catalytically active material and they are designated as inactive only to distinguish themselves from bi-functional catalysts, in which support plays a major role in catalytic function. The present embodiment implicitly assumes the possibility of formation of such synergistic multifunctional combination in certain cases.

Accordingly, purpose of employing a support is strongly reasoned due to numerous factors such as economic, process needs and desired physico-chemical properties. The economic reasons as conceived by inventors are mainly cost reduction by extending accessibility of expensive catalytically active material. Further more process needs as recognized by inventors were sufficient mechanical strength imparted to the catalyst, adjustment of bulk density of formed catalyst, to provide heat sink or heat buffer and to dilute the overactive phase. In addition to these, inventors have recognized geometric needs of catalyst that are primarily satisfied by the support can be described as increased the surface area of the catalyst, optimization of porosity of the overall design. Other chemical features inventor feel necessary to state explicitly in this embodiment are supports provide a means to reduce sintering or deactivation and may also provide acidic or basic centers which function in synergy with catalytically active material.

Although in principle any stable solid material of high surface area, porosity, strength and required texture is suitable, depending on the particular application under consideration. Most stable range of solids employable herein is alumina, silica, magnesia, Titania, zirconia aluminophosphates, charcoal, organic polymers, and compacted clays. These materials are preferred due to their high surface areas, porosity and strength. Apart from these properties they also have low coefficient of thermal expansion.

Nearly all the insulating solids are useful as supports, although on economic grounds alumina and silica are preferred supports. It is recognized from previous reports that oxides such as alumina, silica, zirconia and thoria tend to be acidic. These properties are either of no importance or can be eliminated by selective poisoning. Many naturally occurring materials belong to this group such as pumice asbestos, calcined clays such as bentonite, sepiolite and diatomacious earth such as keisulghur. As a result of wide variation in structure, solids offer range of surface areas and porosity. Although synthetic

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versions of some materials may be preferred in that they offer more closely defined range of properties. In cases where concerted reactions are required wherein one of the reaction may be catalyzed by support it self. The support can catalyze reaction due to acidic and basic sites available on it or metallic sites purposely formed on it. In such cases support by itself is another solid catalyst formed from metal supported on solid support. Illustrative supports belonging to these category of supports are 5% Pd on carbon, 1% Ni on alumina copper-chromite calcined and reduced before use, ruthenium on silica, platinum sulfide on carbon, etc. In considering the individual factors, which dictate the choice of the support, it is realized that the final choice depends on the weighing of these factors in the context of the use to which catalyst are to be employed.

As evident, the rate of a catalytic reaction is dictated by the rate of the chemical reaction on the surface when observed activity is the function of the surface area of the solid support. In practice however overall rate of reaction is usually affected by mass or heat transfer, in which case porosity and geometry of the catalyst particle become increasingly important. As a result choice of support depends on the surface area of the catalyst that can be made available to the reactants and on the porosity of the catalyst.

In context of present invention optimization of surface area is an important factor, which is related to other properties such as texture and the strength. Thus surface area and porosity are closely related, and it can be easily extrapolated further that porosity and mechanical strength is also interrelated. It is obvious to the designer to ensure long life for which catalyst needs a stable structure that is strongly bound together. Certainly this is not the case if porosity is too high. In case of supports of natural origin it is difficult to tailor degree of porosity in systematic fashion. Zeolites or carbon molecular sieves have most of their surface area within the channels, which due to their narrow width restrict passage of reactant molecules. Some gamma aluminas have pore size distribution in the range of 100-200 A⁰, while foamed aluminas have few micropores. Pore diameter can also be increased by careful precipitation of material in pore mouth.

Accordingly in addition to acting as a physical vehicle for the catalytic site support can have appreciable effect on the catalytic reaction it self, wherein for example local pH can be different or bulk of the support can stearically influence course of the reaction and even prevent its occurrence.

As discussed earlier it is understood that apart from chemical behavior of the active phases support plays important role in defining catalyst properties. Such properties could be utilized depending on the process requirements. A considerable advantage would

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be gained if support effects on the active catalytic phases could be minimized which is often difficult in heterogeneous catalysts. By separating effects of active phases and support on can tailor the morphology of the catalyst by selecting support and active phases independent of each other.

Accordingly said support in the catalytic formulation is porous solid pores of which are predominantly of diameter greater than about 20 A⁰ and have a pore diameter in the range of about 3-3000 A⁰. It is also preferred that support material be inert towards substrate, intermediates, products and solvent of the reaction unless concerted tandem sequence of reaction is desired of which one or more reactions is catalyzed by support itself. The suitable catalyst support is any solid that is insoluble in reaction medium and which is thermally stable and high melting solid. The support materials are exemplified but not limited by pumice, alumina-gel, silica gel, silica-alumina-gel, aged or deactivated silica-alumina cracking catalyst, magnesia, diatomaceous earth, bauxite, titania, zirconia, clays, both natural and acid treated, attapulgite, bentonite, diatomaceous earth, keisulghur, lime, calcium carbonate, calcium silicate, magnesium silicate, carborundum, activated and inactivated charcoal, adsorptive carbon, zeolites, zeolite molecular sieves, hydrotalcite, solid foams such as ceramic honeycombs, porous organic polymers such as macroreticular ion exchange resins, poromeric polymers, porous crosslinked polystyrene-sulfonated, calcium alginate, barium sulfate, powdered cellulose, woven cotton mesh, foamed paper, functionalized polymers. It is also possible that the support may be a supported metal catalyst. Above said support materials may be used as regular and irregular particles, capillary tubes, meshes, fabric meshes and inter-spacing elements such as shapes, extrudates, ceramic rods, balls, broken pieces, raschig rings, tiles. Support materials can also have modifiers or deactivators present from impregnation or spraying processes, or other forming operations

As described in the earlier embodiment present invention is concerned with solid phase multi-component formulation in which the catalytically active material is placed on the surface of the solid support. It is invented that soluble catalytic material such as organometallic complexes if rendered insoluble can form said catalytically active solid phases wherein active sites are defined isotropic molecular entities otherwise existing only in solution state. Such insoluble material when dispersed and supported on the surface of the solid support can form simple solid catalyst of the choice. The illustration as depicted in figure 2 outline the strategy as envisaged by the inventors.

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Active components are composed of solid phase that is catalytically active i.e., which is primarily responsible for desired chemical transformation. Unlike support generality of choice for such material is seldom available and composition must be rationally developed within the framework of laws of relevant chemistry. As it is well understood that diversity of materials can catalyze same reaction but one material may not necessarily catalyze diverse range of reactions. It is therefore another specific embodiment of the present invention whereby soluble molecular catalyst are appropriately modified such that they can be incorporated in the said solid material

The desired properties of such catalytically active solid material are,

- 1. Material should not be dissolved or withered in wide variety of reaction medium and conditions
- 2. Said material should have sufficient mechanical and fracture strength
- 3. Such material should be generated from organic organometallic building blocks
- 4. Material should have strong cohesive tendency towards support and total formulation should remain as composite material throughout reaction conditions such as temperatures from -78 to 300 °C, in liquids comprising aqueous, organic and combination thereof as well as in acidic and alkaline conditions
- 5. Said material should be high melting and non subliming
- 6. Catalytically active solid material should be thermally stable should not pyrolyze at reaction temperatures
- 7. One of the building block of the material is molecular component responsible for particular reaction to be catalyzed.

The reaction media as said earlier is quite broad class of liquids and may be selected depending on solubility of substrates and other components as well as it should provide clean recovery of products. The liquids usable as reaction media are exemplified but not limited by petroleum fractions of different boiling ranges, cyclo alkanes such as cyclohexane, cycloheptane, cyclodecane, aromatics such as benzene, toluene, xylenes, ethyl benzene, butylbenzene, alcohols including methanol, ethanol, propanol, butanol, amyl alcohols (linear and branched) higher alcohols, cyclohexanol, phenol, xylinol, cresol, acids such as acetic, propionic, butyric, amides such as formamide, dimethyl formamide, pyrolidone, n methyl pyrolidone, nitriles such as acetonitrile, propinitrile, benzonitrile, esters such as ethylacetate, methylacetate, methyl propionate, methyl

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benzoate methyl propionate, ethers such as diethylether, dibutyl ether, diphenylether, tetrahydrofuran, dioxane, furan, ketones such as acetone, methylethylketone, pentane 2 one, cyclohexanone, nitroaliphatics such as nitromethane, nitroethane, nitropropane, nitroaromatics such as nitrobenzene, 2-nitrotoluene, halogenated solvents such as dichloromethane, chloroform, carbon tetra chloride, 1,2 dichloroethane, chlorobenzene, dichlobenzene other high boiling solvents used for specific purpose include, hexadecane, octadecane, hexatracontane, squalene, chlorinated hydrocarbon oil, liquid paraffin, mineral oil, naphthalene, phenanthrene, methyl naphthalene, high boiling substituted and non substituted organic alcohols, glycol, polyglycols, ethers, polyethers, such as glycerol, polyethyleneglycol, propyleneglycol, erythritol, dulcitol, carbitol, diethyleneglycol, polypropylene glycol, tetraethyleneglycol, 2 ethyl 1 3 hexane diol, 1,5 diethylene glycolmonomethyl methoxypolyethylene glycol, pentanediol, methylbenzylether, 1,2,4-butanetriol, polyphenylether, polybutyleneglycol, bis(phenoxyphenyl)ether, tetraethylene glyco dimethylether, high boiling esters such as diisooctylphthalate, dibutyl phthalate, dioctylphthalate, bis(2-ethylhexyl) phthalate, dinonyl phthalate, butyl benzyl phthalate, bis(2-tetrahydrofurfuryl) phthalate, dipropyl tetrachloro phthalate, dioctyl sebacate, bis(2-ethylhexyl)sebacate, inorganic solvents employable are water, room temperature ionic liquids, flours solvents and super critical dense phases. It is also possible that combination of one or more solvent media be used for reactions depending on solubility of reactants and products. The criteria for selection of solvent are chemical physical requirements of the reaction than the catalyst formulation components. The catalyst formulation as a whole is stable in diverse reaction media so practically any liquid can be used as solvent of the reaction as in case of conventional heterogeneous catalysts. Of course it is understood that for optimum performance of the catalyst very few liquids are suitable and must be selected accordingly.

Properties described earlier for catalytically active solid material are generally found in materials such as ceramics. It is well known in the art that ceramics contain metallic and non-metallic elements that are bonded ionically, covalently or both. These materials can be classified according to their structural composition of which $A_m X_n$ is most common example. A is polyvalent metal cation having +m charge and X is polyvalent anion having -n charge. These materials being ionic lack free electrons making them poor conductor of heat and electricity. Moreover ionic bonds being highly stable and directional also impart high melting range to ceramics. Usually ceramics are also more hard and resistant to physical and chemical changes. Other factors influencing the

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structure and property relationship of ceramic materials can be described as radius ratio and electronegativity difference between positive and negative ions although net negative charge on the material is nil. It was thus clearly envisaged that catalytic material should have properties similar to ceramics

It is thus another embodiment of this work to develop a solid material wherein unit blocks are composed of defined catalytic entities. Another purpose of the present work is that development of a strategy without limiting said catalytic formulation to one particular class of complexes or reaction following certain mechanism according to some particular theory. It was conceived that materials of $A_m X_n$ type if formed where in A_m is a alkaline earth metal cation and X being anion having structure responsible for catalyzing particular reaction then resultant material would have properties similar to ceramics. It was further speculated that such materials being ionic would not be soluble in organic solvent which are customarily used as solvents at the same time such materials due to their negligibly low solubility in aqueous solvents can be employed wherein media is aqueous.

In order to validate and universally substantiate this hypothesis several comparative experiments as referred in examples were undertaken. Anions having two or more negative charges were interacted with group IIA metal cations in solutions. Variety of group IIA compounds including salts, complexes, alkyls and hydrides were interacted with variety of anions having negative charge ranging from -1 to -3 and polyanionic compounds. Various group IIA compounds used for this were selected from magnesium chloride, magnesium acetate, magnesium nitrate, magnesium acac, magnesium complex of ethylene diamine tetraacetic acid disodium salt bytyl magnesium chloride, calcium hydroxide, calcium chloride, calcium nitrate, calcium hydride, calcium acac, calcium complex of ethylene diamine tetra acetic acid disodium salt, strontium acetate, strontium chloride, strontium acac, strontium complex of ethylene diamine tetra acetic acid disodium salt, barium nitrate, barium hydroxide, barium acetate, barium chloride. Such compounds were used as source for group IIA cations in solution. These cations were interacted with anions bearing -1, -2, -3 negative charge and polyanionic compounds. Such anions in solution were obtained from sodium nitrate, sodium propionate, p-toluene sulfonate sodium, m benzene disulfonate disodium, disodium oxalate, disodium sulfate, disodium phenyl phosphonate, disodium hydrogen phosphate, sodium hydrogen phthalate, ammonium molybdate, sodium carboxy methyl cellulose, sodium polyvinyl sulfonate. It was conclusively verified that "group IIA metal ions except magnesium ions form insoluble salts when interacted with anion having at least two or more negative charge.

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Such salts are insoluble in organic, mixture of aqueous organic and have extremely low solubility in water.

Accordingly the hypothesis is confirmed that alkali metal salts containing cation (Aⁿ⁺ wherein n>1) when interacted with polyanions (X), provide a material that is insoluble in majority of solvents including organic as well as aqueous. Such proposition was validated by interacting variety of anionic compounds as described in experiment 1 based on diversity of anion structure it was realized that salts of alkaline earth metal and anion having 2 or more negative charges would provide a solid. Contemplative conclusions were drawn by systematically varying molecular volumes of the polyanions, electron density of the anionic functional groups and alkali metal cations. It was realized that sole requirement for material to form insoluble matter is that anion (X) as described earlier should have at least two anionically charged functional groups. From the experiments detailed subsequently it was realized that anions as small as oxalate to poly anions as large as polyvinyl sulfonate form sparingly to almost insoluble material in water and totally insoluble in organic solvents. If polyanionic nature is introduced on the peripheral positions of the catalyst molecules such that introduction of such groups does not interfere or affect catalytic reaction, would provide anions (X) as said earlier.

Said anionic compounds are those which in conventional sense are acids with proton as counter cation. It is also preferred that strongly acidic functional groups be introduced on the native catalytically active species. Strongly acidic groups are preferred for the reason that these cannot be further protonated in contact with stronger acid that may exist in the reaction medium. It is therefore preferred that strongly acidic functional groups be selected such as for example -SO₃¹⁻, -PO₃²⁻, etc though other groups are also suitable for example -COO provided reaction medium is not acidic.

The catalytically active species described in earlier embodiment is a molecular entity having structural features necessary for intended catalysis. Such molecular entities for example are metal complex catalyst, metal oxoanions or ion pair. Peripheral positions of such catalytically active entities are substituted with anionic functional groups such as SO_3^{1-} , $-PO_3^{2-}$, $-COO^-$ and degree of substitution being essentially > 1.

The substitution/modification as said herein is specifically meant molecular modification of the entity such that it bears said anionic functional groups such as for example -COO⁻, -SO₃⁻, -PO₃²⁻ etc. The term modification doesn't necessarily mean that modified entity is chemically derived from parent entity but it is the analogue of parent

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structure synthesized independently. It is further specified that such anionic functional groups be attached to one of the carbon atoms of the parent entity.

Thus the explicit statement of preferred embodiment is, said catalytic formulation is a combination of solid support as described in earlier embodiment having deposited thereon catalytically active solid material and ensemble as a whole exists as a stable composite solid in gas or liquid phases. The said liquid phases are comprised of aqueous, organic liquids or mixture thereof containing reactants, products and promoters. The catalytic formulation of this embodiment as a physically stable composite solid in gas or liquid phases over a temperature range of -78 to 300 °C and remain as a physically stable composite solid in gas or liquid phases over pressure ranging from 5 to 5000 psi. The catalytically active materials of the embodiment are insoluble salts comprising of group IIA metal, catalytically active inert additive and catalytically active entities. To state further the group IIA metal exists as a cation having +2 charge. The group IIA metal cation of the said catalytically active material is selected from calcium, strontium and barium. And specifically excludes magnesium. The group IIA metal forming catalytically active material may be selected independently or in combination with other group IIA metals. The said catalytic material is formed by precipitation of polyanionic catalytically active entity and catalytically inactive polyanionic entity along with earlier stated group IIA metal ions.

The addition of inert additive is strongly reasoned for reducing solubility of catalytically active solid in aqueous solvent. As said material is ionic tends to dissociate in water and thereby dissolves in liquid phase if it is incidentally liquid. In order to suppress this, other ionic material is required to be additionally present for sacrificial solubility and reduction of solubility by common ion effect, phenomenon that is well known in the literature. Additionally it is also envisaged that addition of such additive provides a microporosity to this material. Addition of catalytically inert material, which by itself is one of the components of the metal ligand complex, provides a surplus coordination capacity to the solid material which acts as significant buffer permitting retention of coordinated transition metal in the complex. Conversely it implies that presence of additional ligand as catalytically active additive prevents the loss of transition metal as well. It is further understood that catalytically inert additive may be present optionally depending on adsorptivity of the support, fluid stresses on the solid particles and coordination tendency of the reactants and solvents. The catalytically active entity or

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entities can be independently selected from metal complexes, quaternary compounds, metal oxo anions and polyoxometalletes or combination thereof.

The metal complexes as described earlier has a general formula

$$(M)_x(L)_y(L^*)_z$$

wherein M is catalytic metal atom or ion of coordination complex is a transition metal from group IIIB, IVB, VB, VIB, VIIB, IB or IIB of the periodic table of elements and may be selected independently suitable transition metal ions and atoms include Sc, Y, Ti, Zr, Hf, V, Cr, Mo, W, Mn, Re, Fe, Ru, Os, Co, Rh, Ir, Ni, Pd, Pt, Cu, Ag, Au, Zn. Preferably the metal complex will contain metal atom or ion from group VIII of periodic table of elements, the suffix x stands for number metal atoms or ions being present from 1 to 60, L is aliphatic, aromatic and heterocyclic compounds containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃,-SO₂, -PO₃², -COO, -O, AsO₃² and -S, the suffix y is required to be at least one. L* is a radical selected from organic anion, inorganic anion and coordinating compound containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene, =C: having optionally attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, aryloxy, cycloalkyl, hydrido, carbonyl, acyl and alkyl and Z is from 0 to 7

As described in earlier embodiment catalytically inert additive is optionally selected from ligand compounds wherein, ligand compounds contain at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, aryloxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃,-SO₂ -PO₃², -COO, -O, AsO₃ and -S

The quaternary compound of as described in one of the previous embodiments has general formula

$[(Y^{+})(R^{*})_{I}][Z^{-}]$

wherein, I = 4 for $Y^+ = N^+$, P^+ , As^+ and I = 3 for $Y^+ = S^+$ and R^* is selected independently from alkyl, aryl, arylalkyl, alkylaryl, alcoxy, aryloxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from $-SO_3^-$, $-SO_2^-$ - PO_3^{2-} , $-COO^-$, $-O^-$, AsO_3^{2-} and $-S^-$ and Z is anion selected from organic anion, inorganic anion or coordination complex anion

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Such type of anionically charged ligands sulfonated tertiary phosphine metal salts ligands employable in this invention and/ or their methods for their manufacture are well known or obvious as seen e.g. by procedures described in "J. Chem. Soc.", pp. 276-288 (1958), US patent nos. 4,483,802 and 4,731,486 for instance such ligands can be prepared by sulfonating corresponding aromatic tertiary phosphine with furning sulfuric acid under controlled temperature conditions to form predominantly protonated di or poly sulfonated phosphines, e.g.

R1,R2 are aryl aryl alky or alkyl

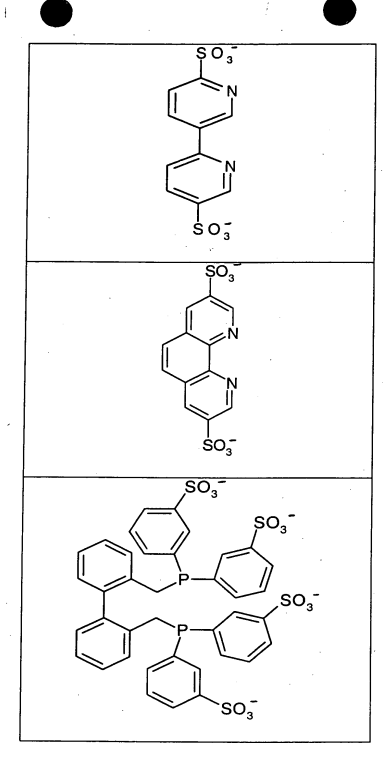
For example the solid phosphine is added to furning sulfuric acid in portions while controlling temperature below 5 °C and then heated, e.g. to 20 –80 °C until desired degree of sulfonation is achieved. The reaction mixture is then cooled immediately to stop any further sulfonation or oxidation of phosphine and without waiting water is added to this avoiding temperature raising above 30 °C and said protonated phosphine salt is neutralized with alkali solution. The mixture containing alkali sulfonate and alkali sulfate is concentrated by evaporating water. During the evaporation of water alkali sulfate precipitate, which is removed by filtration and methanol, was mixed to this mother liquor. Most of the alkali sulfate precipitate and sulfonated phosphine is extracted in the methanol. Evaporation of methanol affords sulfonated phosphine as solid. Dissolving in suitable solvent such as water or ethanol and recrystallizing it therefrom may further purify the crude tertiary phosphine metal sulfonate.

The sulfonation can also be carried out in concentrated sulfuric acid media using boric acid and sulfur trioxide complex as described by Albanese et al US 5684181 and US5780674. The advantage of such procedure is that it reduces phosphine oxide formation. Similarly work up of the sulfonation reaction may also be modified by extracting quenched sulfonation mixture by tributyl phosphite or tri iso octyl amine organic phase which is subsequently extracted with alkali solution advantage of such procedures being phosphines can be selectively separated from corresponding oxides. Oxides of phosphines are frequent contaminants in such sulfonated phosphine ligands.

Presence of phosphine oxide as such doesn't affect catalytic behavior of the ligand in combination with transition metal. It is understood that such phosphine oxides don't coordinate with metal so contamination due to phosphine oxide may be tolerated for the purpose of catalyzing reactions. The presence of phosphine oxides may be intolerant in cases such as bidentate ligands and bidentate chiral ligands. Those experts in the field since can easily realize the situation since phosphine mono oxide of the bidentate ligand will display different coordination behavior. The situation is further complicated while preparing catalysts for enantio selective reactions. In such case phosphine oxide removal is desired and may be achieved by extractive separation from tributyl phosphite or tri iso octylamine solutions or by fractionating from gels of modified dextran such as SEPHADEX G15 (TM) as described by Hermann et al Angew. Chem. Int. Eng. Ed. 29 (1990) No (4) 391-393.

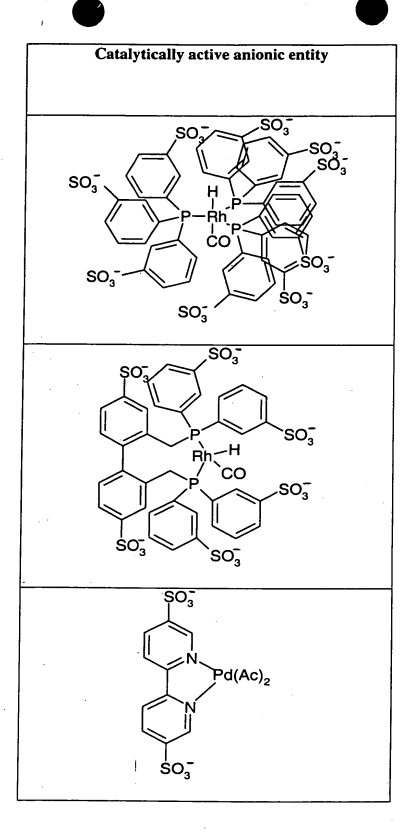
Such ligands that are sulfonated can be prepared by various methods employing lithium phosphides, Grignard reagent and phosphorus trichloride etc. Knowledge and understanding of such ligands is taught in literature known to artisan skilled in the art. For example Kosolapoft G. M., Maier L. Organic Phosphorus Compounds, Volume 1, 288, Wiley Interscience, New York, © 1972./Engl R. synthesis of carbon phosphorus bonds, © CRC Press 1988./ Tripett S. A. A specialist Periodical Report of Organophosphorus Chemistry, Chemical Society London © 1970/ specific example of such synthesis is explained in (Mann, F. G. et al, J. Chem. Soc. 1937, 527-535; US 4,483,802 and US 4,731,486) Similarly nitrogen containing ligands can be prepared by specific chemical synthesis known in the art (Eit Drent, US 5,166,411). Synthesis, manufacture and purification of such ligands is clearly out of purview of this application. It is clearly understood knowledge concerning sulfonation of such ligands is also well known. Similarly ligands bearing other anionically charged functional groups for example COO, -PO3 ²⁻ can also be prepared by sequence of specific organic synthesis (synthesis of phosphonic and carboxylic containing ligands). This invention claims

further utilization of such well known ligands for preparation of generic catalyst formulation which is a solid and employed as a catalyst. This utilization of such known ligands by further processing is within the scope and purview of this application and is one of the preferred embodiments of this invention. Illustrative preferred anionic ligands and their transition metal complexes and quaternary compounds include, as follows. It is implicit that these are only illustrative and not comprehensive



Some of the illustrative quaternary compounds are

Poly anionic ligands and quaternary compounds as described above exist as salts of alkali metal, quaternary ammonium or proton. It is well known in the art that such compounds are water soluble combination of such ligands with transition metals provide access to their complexes. Some of illustrative catalytically active entities are:



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Complexes as said earlier can be prepared by various methods known in literature descriptions. Broadly such metal complexes can be classified as follows:

- 1. Synthesis of metal complexes from metal salts and anionic ligands
- 2. Displacement of a labile ligand by anionic ligand

For example complexes such as PdCl₂ bis (triphenyl phosphine trisulfonate trisodium) is prepared by reacting PdCl₄² with triphenyl triphenyl phosphine trisulfonate trisodium in aqueous ethanol, ruthenium chloride with triphenyl triphenyl phosphine trisulfonate trisodium, where as complexes such as HRhCO(TPPTS)₃, RuCl₂ BINAP, are prepared by displacement of labile ligands such as cyclooctadiene or acetylacetonate. Complexes such as sulfonated pthalocynine are prepared by simultaneous formation of ligand and complex. As said earlier synthesis of such complexes is a well-known knowledge to those experts in the art. The synthesis of such complexes is beyond the scope and purview of this invention. Yet utilization of such complexes to form insoluble solid formulation for catalytic application is the explicit embodiment of this invention.

Several different metal complexes containing different metals and diversity of ligands were synthesized. Such metal complexes whenever interacted with group IIA metal compounds except magnesium compounds provided solids, which were insoluble in water.

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organic solvents. The precipitates were solids up to 200°C and were non-subliming. Several experiments as described in examples were carried out to verify proposed hypothesis to deduce a logical conclusion that when poly-anions when interacted with a group IIA metal, form a precipitate that is insoluble in variety of liquids. It is yet another preferred embodiment of this application that the admixture of catalytically active poly anion and catalytically inactive anion is preferred to form a precipitate that is insoluble in majority of liquids. Additional presence of catalytically inactive additive is reasoned to reduce solubility of precipitated complex in water by well-known phenomena of common ion effect. It is also preferred to add additional ligand that is used to form complex. The presence of additional ligand is preferred especially in cases where catalytic formulation is intended to be used in liquid phases that are coordinating or ligand involved is monodentate.

The catalytically active material of the said catalyst formulation is formed from interaction of solution of catalytically inactive additive, catalytically active entity and a solution of group IIA metal cation by precipitation. The solution of catalytically inactive additive, catalytically active entity when contacted with a solution of group IIA metal cation, two solution start diffusing and subsequently whenever cation of group IIA metal encounters collision with poly anion solidification is initiated and cluster of solid is slowly formed. Such precipitation wherein, catalytically inactive additive is independently selected from anions having at least two or more negative charges, ligand compounds containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, aryloxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃-SO₂-PO₃²-COO-, -O-, AsO₃² and -S-, and combination thereof and catalytically active entity is independently selected from metal complexes, quaternary compounds, metal oxo anions and polyoxometallates or combinations thereof

The catalytically active may be selected such that metal complexes has a general formula

$(M)_x(L)_y(L^*)_z$

wherein M is catalytic metal atom or ion of coordination complex is a transition metal from group IIIB, IVB, VB, VIB, VIB, IB or IIB of the periodic table of elements and may be selected independently suitable transition metal ions and atoms include Sc, Y, Ti, Zr, Hf, V, Cr, Mo, W, Mn, Re, Fe, Ru, Os, Co, Rh, Ir, Ni, Pd, Pt, Cu, Ag, Au, Zn. Preferably the metal complex will contain metal atom or ion from group VIII of periodic table of elements, x is from 1 to 60, L is aliphatic, aromatic and heterocyclic compounds containing

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at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃⁻,-SO₂⁻-PO₃²-, -COO⁻, -O⁻, AsO₃²- and -S⁻, y is at least 1, L* is a radical selected from organic anion, inorganic anion and coordinating compound containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene, =C: having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl, z is from 0 to 7 and quaternary ammonium compound has a general formula

 $[(Y^{+})(R^{*})_{I}][Z^{-}]$

wherein, I = 4 for $Y^{+} = N^{+}$, P^{+} , As^{+}

I = 3 for Y⁺ = S⁺ and R^{*} is selected independently from alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from $-SO_3^-$, $-SO_2^ -PO_3^{2-}$, $-COO^-$, $-O^-$, AsO_3^{2-} and $-S^-$ and Z is anion selected from organic anion, inorganic anion or coordination complex anion.

Accordingly as described in earlier embodiment inventors have developed a common technique for solidification of pluralistic catalytic entities, functioning according to different theories and mechanisms. Such solidification is achieved by incorporating earlier said catalytic entities in ionic solid, which by itself is formed by interaction of poly anionic catalytic entity, polyanionic additive and group IIA metal ion.

The catalytic formulation of this invention wherein earlier said catalytic material is supported on the surface of the solid support. It is recalled that formed catalytic material being insoluble cannot be dissolved in liquid and then supported on the solid support accordingly technique was required to form such material directly on the surface of the solid support by precipitation.

The final formation of composite catalyst can thus be carried out by precipitation of catalytically active material on the support surface. The formation of catalyst by precipitation or co precipitation is thus centrally important in this respect. However precipitation is a complex phenomenon and demand several ancillary techniques to be developed in order to deposit catalyst on the support surface. Nevertheless, for several catalytically relevant materials especially for support materials precipitation is most frequently applied method. In this respect such precipitation is troublesome as it may generate clusters and particles in the bulk of liquid. Dealing specifically formation of solid catalytically active material is better described by term co precipitation as two components categorized as group IIA metal ion and poly anionic entity when interacted yields a precipitate. Co precipitation is extremely suitable technique for generation of uniform

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distribution of such material on the support material, as stoichometry of interacting species is definite. Form earlier experiences it is known fact that co precipitation can provide good dispersion of the support surface which is otherwise difficult to achieve catalyst assembly that is under consideration. Thus the bulk co precipitation process needs to be modified to achieve assembly of composite catalyst system.

Preferably, the co precipitation is carried out in such a manner that precursor solutions containing anionic entities (catalytically active entity and catalytically inert additive) and group IIA salt solutions diffuse near the surface of the support or the formation of insoluble clusters initiate near the surface of the support. Hereinafter solution containing anionic component is designated as solution A and solution containing group IIA metal is designated as solution B. It is another embodiment of the present invention that outlines various methods for assembling catalytic formulation of earlier said embodiments. These assembly techniques are broadly classified according to various techniques of precipitation and are described in ensuing description of embodiments.

One of the process for the preparation of a heterogeneous catalytic formulation as a solid composite comprising of porous solid support having deposited thereon a catalytically active solid is characterized by suspending insoluble solid support in a liquid phase to which a solution of catalytically inert additive and catalytically active entity and a solution of group IIA metal cation are added simultaneously or sequentially with vigorous agitation and allowed to age for 1 to 48 hours wherein, support is a mechanically robust and thermally stable solid in reaction media, having a mean pore diameter in the range of about 3-3000 A⁰ and existing as powder, granules, flakes or pallets of regular or irregular shapes, sheets, monolith, ropes and woven fabric of fibrous solids and catalytically inactive additive is independently selected from anions having at least two or more negative charges which may be organic, inorganic, or a compound containing at least one radical form O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃,-SO₂ -PO₃²⁻, -COO, -O, AsO₃ and -S.

The catalytically active entity is independently selected from metal complexes, quaternary compounds, metal oxo anions and polyoxometallates or combinations thereof such that metal complexes having a general formula

$$(M)_x(L)_y(L^*)_z$$

wherein M is catalytic metal atom or ion of coordination complex is a transition metal from group IIIB, IVB, VB, VIB, VIIB, IB or IIB of the periodic table of elements and may

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be selected independently suitable transition metal ions and atoms include Sc, Y, Ti, Zr, Hf, V, Cr, Mo, W, Mn, Re, Fe, Ru, Os, Co, Rh, Ir, Ni, Pd, Pt, Cu, Ag, Au, Zn. Preferably the metal complex will contain metal atom or ion from group VIII of periodic table of elements, x is from 1 to 60, L is aliphatic, aromatic and heterocyclic compounds containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃,-SO₂ -PO₃²⁻, -COO⁻, -O⁻, AsO₃²⁻ and -S⁻ and y is at least 1 and L* is a radical selected from organic anion, inorganic anion and coordinating compound containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene, =C: having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl, z is from 0 to 7 and the quaternary ammonium compound has a general formula

$[(Y^{+})(R^{*})_{I}][Z^{-}]$

wherein, I = 4 for $Y^+ = N^+$, P^+ , As^+ , I = 3 for $Y^+ = S^+$ and R^* is selected independently from alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from $-SO_3^-$, $-SO_2^-$, $-PO_3^{2-}$, $-COO^-$, $-O^-$, AsO_3^{2-} and $-S^-$ and Z is anion selected from organic anion, inorganic anion or coordination complex anion. The group IIA metal cation is selected from compounds of Ca^{2+} , Sr^{2+} and Ba^{2+} .

The above said process is carried out in the temperature ranging from -78 to 200 °C preferably between -5 to 100 °C. The solvent for the process is selected from aqueous, water miscible organic or mixture thereof.

The process as described above wherein solution of catalytically inert additive and catalytically active entity and a solution of group IIA metal cation are added simultaneously over a period of 10 to 1500 min. After completion of this treatment the catalyst is recovered by centrifugation, decantation, gravity settling or other techniques of solid liquid separation and dried subsequently in vacuum. The method as described herein is employable when components of precipitate slowly produce solid material under the influence of viscosity, solvent media and solubility modifiers. As seeds of solid material develop slowly and there is enough time for seeds of the precipitate to settle on the support surface. Other methods described herein after are suitable for co-precipitation that occurs instantaneously. Such methods are usually critical due to specialized unit operation required for them and also require specific equipment for the manufacture.

Another process for the preparation of a heterogeneous catalytic formulation as a solid composite comprising of porous solid support having deposited thereon a

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catalytically active solid is characterized by impregnating the solid support with catalytically active entity and catalytically inert additive followed by drying, dried support having deposited thereon catalytically active entity and catalytically inert additive is added to a solution of group IIA metal compound, with simultaneous agitation. The suspension is aged for 1 to 48 hours with agitation, the process is accordingly carried out in the temperatures ranging from -78 to 200 °C preferably between -5 to 100 °C

The support in this case is a mechanically robust and thermally stable solid in reaction media, having a mean pore diameter in the range of about 3-3000 A⁰ and existing as powder, granules, flakes or pallets of regular or irregular shapes, sheets, monolith, ropes and woven fabric of fibrous solids and catalytically inactive additive is independently selected from anions having at least two or more negative charges which may be organic, inorganic, or a compound containing at least one radical form O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃,-SO₂ -PO₃², -COO⁻, -O⁻, AsO₃² and -S⁻

The catalytically active entity is independently selected from metal complexes, quaternary compounds, metal oxo anions and polyoxometallates or combinations thereof such that metal complexes having a general formula

$$(M)_x(L)_v(L^*)_z$$

wherein M is catalytic metal atom or ion of coordination complex is a transition metal from group IIIB, IVB, VB, VIB, VIIB, IB or IIB of the periodic table of elements and may be selected independently suitable transition metal ions and atoms include Sc, Y, Ti, Zr, Hf, V, Cr, Mo, W, Mn, Re, Fe, Ru, Os, Co, Rh, Ir, Ni, Pd, Pt, Cu, Ag, Au, Zn. Preferably the metal complex will contain metal atom or ion from group VIII of periodic table of elements, x is from 1 to 60, L is aliphatic, aromatic and heterocyclic compounds containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃,-SO₂ -PO₃², -COO, -O, AsO₃² and -S and y is at least 1 and L* is a radical selected from organic anion, inorganic anion and coordinating compound containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene, =C: having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl, z is from 0 to 7 and quaternary ammonium compound has a general formula

 $[(Y^{+})(R^{*})_{I}][Z^{-}]$

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wherein, I = 4 for $Y^+ = N^+$, P^+ , As^+ ; I = 3 for $Y^+ = S^+$ and R^* is selected independently from alkyl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from $-SO_3$, $-SO_2$, $-PO_3^{-2}$, $-COO^-$, $-O^-$, AsO_3^{-2} and $-S^-$ and Z is anion selected from organic anion, inorganic anion or coordination complex anion. The group IIA metal cation is selected from compounds Ca^{2+} , Sr^{2+} and Ba^{2+} . According to process under consideration solvents used to dissolve anionic components and group IIA metal cations are aqueous, water miscible organic or mixture thereof

The process modification may be adopted wherein support having deposited thereon catalytically active entity and catalytically inert additive is added to a solution of group IIA metal compound, with simultaneous agitation over a period of 10 to 1500 min, depending upon specific process requirements. The process accordingly concludes by recovering catalyst by centrifugation, decantation, gravity settling or other techniques of solid liquid separation and drying subsequently in vacuum

Yet according to another preferred process for the preparation of a heterogeneous catalytic formulation as a solid composite comprising of porous solid support having deposited thereon a catalytically active solid is characterized by impregnation of support with a solution of catalytically inactive additive and catalytically active entity followed by drying. Solid support having deposited thereon catalytically inactive additive and catalytically active entity is suspended in water immiscible solvent to which a solution of group IIA metal compound is added with vigorous agitation and concurrent removal of low boiling or azeotropic fraction of a solvent. Suspension is allowed to age for 1 to 48 hours, wherein the process is accordingly carried out in the temperature ranging from 70 to 200°C

The support in this case is a mechanically robust and thermally stable solid in reaction media, having a mean pore diameter in the range of about 3-3000 A⁰ and existing as powder, granules, flakes or pallets of regular or irregular shapes, sheets, monolith, ropes and woven fabric of fibrous solids and catalytically inactive additive is independently selected from anions having at least two or more negative charges which may be organic, inorganic, or a compound containing at least one radical form O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃,-SO₂ -PO₃², -COO, -O, AsO₃² and -S

The catalytically active entity is independently selected from metal complexes, quaternary compounds, metal oxo anions and polyoxometallates or combinations thereof such that metal complexes having a general formula

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$(M)_x(L)_y(L^*)_z$

wherein M is catalytic metal atom or ion of coordination complex selected from a transition metal groups IIIB, IVB, VB, VIB, VIIB, IB or IIB of the periodic table of elements and may be selected independently suitable transition metal ions and atoms include Sc, Y, Ti, Zr, Hf, V, Cr, Mo, W, Mn, Re, Fe, Ru, Os, Co, Rh, Ir, Ni, Pd, Pt, Cu, Ag, Au, Zn. Preferably the metal complex contains metal atom or ion from group VIII of periodic table of elements, x is ranging from 1 to 60, L is selected from aliphatic, aromatic and heterocyclic compounds containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃,-SO₂ -PO₃², -COO, -O, AsO₃² and -S and y is at least 1 and L* is a radical selected from organic anion, inorganic anion and coordinating compound containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene, =C: having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl, z is from 0 to 7 and the quaternary ammonium compound has a general formula

$[(Y^{+})(R^{*})_{I}][Z^{-}]$

wherein, I = 4 for $Y^+ = N^+$, P^+ , As^+ , I = 3 for $Y^+ = S^+$ and R^* is selected independently from alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃, -SO₂, -PO₃², -COO, -O, AsO₃² and -S and Z is anion selected from organic anion, inorganic anion or coordination complex anion. The group IIA metal cation is selected from compounds of Ca^{2+} , Sr^{2+} and Ba^{2+} .

The solvents employed to form a solution of group IIA metal ion are aqueous, water miscible organic or mixture thereof and solvent employed to suspend support is water immiscible organic solvent having boiling point in the range 40 to 200 °C, the present process is concludes by recovering catalyst by centrifugation, decantation, gravity settling or other techniques of solid liquid separation and dried subsequently in vacuum

Process for the preparation of a heterogeneous catalytic formulation as a solid composite comprising of porous solid support having deposited thereon a group IIA metal compound followed by drying. Solid support having deposited thereon group IIA metal is suspended in water immiscible solvent to which a solution of catalytically active entity and catalytically inactive additive is added with vigorous agitation and concurrent removal of low boiling or azeotropic fraction of solvent. The process of azeotropic distillation is accordingly carried out in the temperature ranging from 70 to 200 °C. The liquid medium

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employed for the process of azeotropic removal of solvent is water immiscible organic solvent having boiling point in the range 40 to 200 °C. There after suspension is allowed to age for 1 to 48 hours.

The support in this case is a mechanically robust and thermally stable solid in reaction media, having a pore diameter in the range of about 3-3000 A⁰ and existing as powder, granules, flakes or pallets of regular or irregular shapes, sheets, monolith, ropes and woven fabric of fibrous solids and catalytically inactive additive is independently selected from anions having at least two or more negative charges which may be organic, inorganic, or a compound containing at least one radical form O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃, -SO₂-PO₃², -COO⁻, -O⁻, AsO₃² and -S⁻

The catalytically active entity is independently selected from metal complexes, quaternary compounds, metal oxo anions and polyoxometallates or combinations thereof such that metal complexes has a general formula

$$(M)_x(L)_y(L^*)_z$$

wherein M is catalytic metal atom or ion of coordination complex selected from transition metal group IIIB, IVB, VB, VIB, VIIB, IB or IIB of the periodic table of elements and is selected independently suitable transition metal ions and atoms include Sc, Y, Ti, Zr, Hf, V, Cr, Mo, W, Mn, Re, Fe, Ru, Os, Co, Rh, Ir, Ni, Pd, Pt, Cu, Ag, Au, Zn. Preferably the metal complex will contain metal atom or ion from group VIII of periodic table of elements, x is from 1 to 60, L is aliphatic, aromatic and heterocyclic compounds containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃,-SO₂ -PO₃², -COO, -O, AsO₃² and -S and y is at least 1 and L* is a radical selected from organic anion, inorganic anion and coordinating compound containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene, =C: having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl, z is from 0 to 7 and the quaternary ammonium compound has a general formula

$$[(Y^{+})(R^{*})_{I}][Z^{-}]$$

wherein, I = 4 for $Y^+ = N^+$, P^+ , As^+ , I = 3 for $Y^+ = S^+$ and R^+ is selected independently from alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from $-SO_3^-$, $-SO_2^ -PO_3^{2-}$, $-COO_3^-$, $-O_3^-$, AsO_3^{2-} and $-S_3^-$ and Z is anion selected from organic anion, inorganic anion or

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coordination complex anion. The group IIA metal cation is selected from compounds of Ca²⁺, Sr²⁺ and Ba²⁺. And solvent employed to form a solution of group IIA metal ion is aqueous, water miscible organic or mixture thereof. After removal of organic immiscible and low boiling liquids, centrifugation, decantation, gravity settling or other techniques of solid liquid separation and dried subsequently in vacuum to recover the catalyst

Process for the preparation of a heterogeneous catalytic formulation as a solid composite comprising of fluidizing solid support in the current of gasses. Solution of catalytically active entity and catalytically inert additive is sprayed in such a way that catalytically active entity and catalytically inert additive is deposited on the solid support the fluidization of solid is continued for 1 to 48 hours. Solution of group IIA metal compound is subsequently sprayed and fluidization of solid is further continued for 1 to 48 hours and solids are recovered. The process of fluidization is carried out in the temperature ranging from 20 to 200 °C, wherein the support in this case is a mechanically robust and thermally stable solid in reaction media, having a mean pore diameter in the range of about 3-3000 A⁰ and existing as powder, granules, flakes or pallets of regular or irregular shapes, sheets, monolith, ropes and woven fabric of fibrous solids and catalytically inactive additive is independently selected from anions having at least two or more negative charges which may be organic, inorganic, or a compound containing at least one radical form O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃, -COO, -O, AsO₃² and -S the catalytically active entity is independently selected from metal complexes, quaternary compounds, metal oxo anions and polyoxometallates or combinations thereof such that metal complexes has a general formula

 $(M)_x(L)_y(L^*)_z$

wherein M is catalytic metal atom or ion of coordination complex is a transition metal from group IIIB, IVB, VB, VIB, VIIB, IB or IIB of the periodic table of elements and may be selected independently suitable transition metal ions and atoms include Sc, Y, Ti, Zr, Hf, V, Cr, Mo, W, Mn, Re, Fe, Ru, Os, Co, Rh, Ir, Ni, Pd, Pt, Cu, Ag, Au, Zn. Preferably the metal complex will contain metal atom or ion from group VIII of periodic table of elements, x is from 1 to 60, L is aliphatic, aromatic and heterocyclic compounds containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃,-SO₂

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-PO₃²⁻, -COO⁻, -O⁻, AsO₃²⁻ and -S⁻ and y is at least 1 and L^{*} is radical selected from organic anion, inorganic anion and coordinating compound containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene, =C: having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl, z is from 0 to 7 and quaternary ammonium compound has a general formula

$[(Y^{+})(R^{*})_{I}][Z]$

wherein, I = 4 for $Y^+ = N^+$, P^+ , As^+ , I = 3 for $Y^+ = S^+$ and R^* is selected independently from alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃, -SO₂ -PO₃²⁻, -COO⁻, -O⁻, AsO_3^{2-} and -S⁻ and Z is anion selected from organic anion, inorganic anion or coordination complex anion. The group IIA metal cation are selected from compounds of Ca^{2+} , Sr^{2+} and Ba^{2+} . The solvent employed to form a solution of group IIA metal ion is aqueous, water miscible organic or mixture thereof.

This invention further extends another preferred method for making catalysts according to present invention. According to this method an anionically charged entity and anionically charged additive are deposited on the solid support and are subsequently cured by spraying group IIA metal salt solution with simultaneous removal of solvent.

Accordingly, process for the preparation of a heterogeneous catalytic formulation as a solid composite comprises of tumbling solid support in the rotating pan under current of inert gasses. Solution of catalytically active entity and catalytically inert additive is sprayed in such a way that catalytically active entity and catalytically inert additive is uniformly deposited on the solid support the tumbling of solid is continued for 1 to 48 hours. Solution of group IIA metal compound is subsequently sprayed and tumbling of wet solid is further continued for 1 to 48 hours and solids are recovered. The process described accordingly is carried out in the temperature ranging from 20 to 200 °C. Either heating the inert gas stream or rotating pan, which contains support, may achieve the process temperature. The laboratory apparatus employed to form present formulation is represented in figure 6 and such apparatus may be suitably scaled depending upon volume requirements.

The support material employable herein is a mechanically robust and thermally stable solid in reaction media, having a mean pore diameter in the range of about 3-3000 A⁰ and existing as powder, granules, flakes or pallets of regular or irregular shapes, sheets, monolith, ropes and woven fabric of fibrous solids.

The catalytically inactive additive is independently selected from anions having at

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least two or more negative charges which may be organic, inorganic, or a compound containing at least one radical form O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from $-SO_3^-, -SO_2^- -PO_3^{2-}$, $-COO^-$, $-O^-$, AsO_3^{2-} and $-S^-$. Additionally this catalytically inactive additive may be polymer bearing multiple anionic charges.

Catalytically active entity is independently selected from metal complexes, quaternary compounds, metal oxo anions and polyoxometallates or combinations thereof. The metal complex entity that is catalytically active can be selected such that metal complexes have a general formula

$(M)_x(L)_y(L^*)_z$

wherein M is catalytic metal atom or ion of coordination complex is a transition metal from group IIIB, IVB, VB, VIB, VIIB, IB or IIB of the periodic table of elements and may be selected independently suitable transition metal ions and atoms include Sc, Y, Ti, Zr, Hf, V, Cr, Mo, W, Mn, Re, Fe, Ru, Os, Co, Rh, Ir, Ni, Pd, Pt, Cu, Ag, Au, Zn. Preferably the metal complex will contain metal atom or ion from group VIII of periodic table of element. The suffix x indicates number of such catalytic transition metal present in the complex. The number of such metal entities ranges from 1 to 60. The component L of the metal complex is aliphatic, aromatic and heterocyclic compounds containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from -SO₃-, -SO₂-PO₃²-, -COO. -O. AsO₃² and -S. The suffix y indicate the number of coordinating ligands that hold metal in the lower oxidation state and it is necessary that y is at least 1. L* is a radical selected from organic anion, inorganic anion and coordinating compound containing at least one radical from O, N, S, Se, Te, P, As, Sb, Bi, Si, olefin, carbene, =C: having attached thereto oxy, alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl. Suffix z indicates number of such non-participating ligands. These ligands may be identical if present in multiple or different but total number ranges from 0 to 7. Alternatively another class of catalytically active entity, which is employable alone or in combination with above said transition metal complex, is quaternary compound which, has a general formula

$[(Y^{+})(R^{+})_{I}][Z^{-}]$

wherein, compounds that are elected can belong to quaternized compounds of nitrogen phosphorus, arsenic and sulfur. Instances when quaternary compound belongs to nitrogen,

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phosphorus, arsenic containing compounds suffix I = 4 for $Y^+ = N^+$, P^+ , and for sulfur compounds I = 3 for $Y^+ = S^+$ R^+ is selected independently from alkyl, aryl, arylalkyl, alkylaryl, alcoxy, arlyoxy, cycloalkyl bearing at least one or more negatively charged functional groups independently selected from $-SO_3^-, -SO_2^- -PO_3^{2-}$, $-COO^-, -O^-$, AsO_3^{2-} and $-S^-$, Z is anion selected from organic anion, inorganic anion or coordination complex anion. In majority cases it is implicit that actual catalytic entity is anion Z Quaternary compound provides an anchor for solidification as well as for providing required electrostatic field such that anion z- does not get away

The group IIA metal cation is selected from compounds of Ca²⁺, Sr²⁺ and Ba²⁺. The process for formation of catalyst in coating pan as described earlier wherein the solvent employed to form solutions is preferably aqueous, water miscible organic or mixture thereof. Such solutions according to process are sprayed simultaniously or sequentially

Irrespective of the processes employed to form catalytic formulation said solid catalytic ensemble could be employed to catalyze diversity of reactions in gas phase or in liquid slurry. The catalyst being robust solid provides an opportunity to select suitable reactor configuration for manufacture of organic compounds in variety of reactor configurations such as fixed bed, trickel bed, fluidized bed and slurry reactors depending on the physical state and properties of reactants and products.

The solid catalyst of the present invention can be optionally modified wherein a film of high boiling liquid or low melting solid is optionally deposited on the solid catalyst. This modification can be adopted to enhance local solubility of reactants or modify environment of the catalytic sites to obtain high selectivity for required products.

The catalyst to be formulated according to earlier described embodiments for particular reaction is selected from analogous catalysts that catalyze such reaction in liquid phase; analogues entity is derived from such parent catalyst of homogeneous system by appropriate functionalization so as to introduce negative charges on it. Catalytically active entity is independently derived from metal complexes, quaternary compounds, metal oxo anions and polyoxometallates or combinations thereof depending upon requirement. Except anionic functional groups rest structure of catalytic entity in immaterial for solidification of such entity. Some of illustrative derivations of anionically charged entities from respective soluble catalysts are displayed in following table. Reaction classes represented are exemplary only and catalytic entities are within the purview of appended claims. It is thus explicit clarification of the embodiment that solid catalyst formulation is described wherein the catalytic entities are solidified by generic technique irrespective of

reaction they catalyze. Such catalytic entities are clearly claimed in claims.

Some of the illustrative examples of anionically functionalized soluble catalytic entities and their applications are listed in following table.

Jr 9/3/03

Reaction type	Soluble catalyst	Analogus anionic entity	
Hydroformylation	HRhCO(TPP) ₃	HRhCO(TPPTS) ₃	
	HRhCO(BISBI) ₃	HRhCO(BISBIS) ₃	
	$Co_2CO_4[P(_nBu)_3]_2$	Co ₂ CO ₄ [TPPTS] ₂	
	SnCl ₃ PtCl (TPP) ₂	SnCl ₃ PtCl (TPPTS) ₂	
Hydrogenation			
	RhCl(TPP) ₃	RhCl(TPPTs) ₃	
	$RuCl_2(C_6H_5)BINAP(S)$	RuCl ₂ (C ₆ H ₅)BINAP(S)	
	RhClO ₄ Chiraphos (S,S)	RhClO ₄ Chiraphossulfonate d (S,S)	
Carbonylation			
	PdAcPTSA(TPP) ₂	PdAcPTSA(TPP) ₂	
	$Rh(CO)_2I_2$ [MeN ⁺ (Ph) ₃]	Rh(CO) ₂ I ₂ [MeN ⁺ (PhmSO ₃	
Heck olefination			
	PdCl ₂ (PPh ₃) ₂	PdCl ₂ (TPPTS) ₂	
Suzuki coupling			
	PdCl ₂ (PPh ₃) ₂	PdCl ₂ (TPPTS) ₂	
Isomarization			
	PtCl ₂ (PPh ₃) ₂	PtCl ₂ (TPPTS) ₂	
	RhCl(TPP) ₃	RhCl(TPPTs) ₃	
Wacker oxidation			
	Pd(Ac) ₂ BIPY	Pd(Ac) ₂ BIPYDS	
Oxidation			
	CoPthalocynine	CoPthalocyninetetrasulfonat ed	
Michel and Knovengel reactions			
	NaOH	OH [MeN (PhmSO ₃) ₃]	
	NaOMe	$MeO^{-}[MeN^{+}(PhmSO_{3}^{-})_{3}]$	

The supported catalysts according to the invention are extremely active as is born out by the tests in the description, which is given in the examples, which follow. In fact these examples relate to the application of such supported catalysts to the diversity of reactions catalyzed by different mechanisms and according to known theories of molecular catalysts. A comparison of the reactions of these supported catalysts in homogeneous

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phase establishes that while retaining catalyst activity to considerable extent facile separation can be easily achieved. This makes catalyst suitable for continuous process thereby enhancing catalyst process economics. The said catalyst formulation being inherently solid can be easily recovered after the desired catalytic conversion in the heterogeneous phase. They can then be reused to catalyze new charge of reactants, this operation being either continuos or repeatable wherein the catalyst can be recycled for several times. The advantageous fact is catalytic formulation being repeatable several times without their activity being appreciably degenerating.

Before subjecting catalysts for purpose of catalyzing reactions it is essential that stability and incompatibilities be assessed. For this reason various chemical stresses were applied to simulate stresses encountered by the catalytic formulation when they are applied to actual reaction. The stresses encountered during reaction or during post processing are solvation stresses due to solvents and media. Ranges of solvents that are applicable are liquids employed for reaction and post processing such as washing. Washing is preferred process to regenerate catalyst to remove adsorbed material and for activation by other chemical treatment.

For this reason various catalysts containing different metals such as rhodium, ruthenium, iridium, palladium, platinum, cobalt, nickel, molybdenum and iron were prepared according to methods described earlier and extracted at boiling temperatures of the solvents like, water, acetic acid, methanol, isopropanol, ether, tetrahydrofuran, ethyl acetate, acetone, acetonitrile, toluene, cyclohexane in the apparatus detailed in figure 3. The extraction was continued for several hours and subsequently solvent was changed. No appreciable loss of metal content was and physical morphology by visual comparison was detected. It is obviously true that sites containing hydroxy, methoxy and other basic radicals would be destroyed. These experiments indicate that catalyst would be stable in diverse range of solvents and need not be restricted to particular class of solvents. Similarly catalysts were leached in aqueous acids and alkaline solutions and loss of metals including transition metal or group IIA metal was detected

The activity of the catalyst studied has been measured in the examples in conventional manner by the turnover number which defines number of molecules converted by the catalytic reaction for a catalytically active entity per unit time under idealized conditions or as yield over a defined period of time.

Various exploratory experiments were carried out to ascertain applicability of the catalyst of the invention. The exploratory experiments were aimed at understanding

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molecular catalysis. Due to this reason reaction was selected where two products are formed with dissymetric regioselectivity. Hydroformylation is one such reaction wherein; reaction rates and regioselectivities are significantly altered because of variation of molecular environment. Due to these reasons hydroformylation of hexene with HRhCO(TPPTS)₃ was considered as suitable probe to understand catalysis in reaction conditions.

Hydroformylation reaction was carried out for hexene as substrate and HRhCO(TPPTS)₃ as active catalytic entity (g of rhodium/g of support), moisture content (ppm), silica as support and no excess of ligand. Total conversion was obtained and the catalyst was recovered by centrifugation, washed with toluene, and dried in vacuum. Dry catalyst powder was reused for hydroformylation of allyl alcohol in water catalyst was active for hydroformylation providing aldehydes. Catalyst after reaction was recovered by centrifugation, dried in vacuum, reused for hexene hydroformylation, and found to produce aldehydes. This experiment indicated feasibility of catalyzing reaction in various solvents successively irrespective of substrate.

Accordingly need of support was identified by precipitating HRhCO(TPPTS)₃ with barium nitrate precipitate was used to catalyze hexene hydroformylation. After 24 hours, conversion was below 1%.

In order to identify that reactions take place in solid state and not by leaching of complex under reaction conditions, which eventually return to solid. This is verified by using criteria of mobility of catalyst species and additional ligand. In case of soluble catalyst wherein additional ligand is in mobile condition due to which it can interact with active species and there by giving lower rates and high n/I ratio. When catalysts were prepared with additional ligand present no change in activity and selectivity was observed. This observation was attributed to immobile state of ligands and catalyst. Due to immobility of ligands their interaction with active species is totally retarded thereby not affecting rates and n/i ratio.

Immobility of catalyst was further verified by addition of water to the solid catalyst. At lower water content (ppm/g) high conversions were obtained. When moisture content was increased activity was considerably reduced. Same catalyst when dried resumed its original activity. This experiment conclusively verifies that reaction occurs in solid state.

Accordingly the crucial evaluation indicating life of the catalyst, its stability and the durability was performed in a tubular fixed bed reactor by subjecting catalyst to

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hydroformylation in tubular trical bed reactor (ϕ ½") at 80 0 C and 300 psi H₂/CO (1:1) using 5 g. of catalyst. 5 % decene in toluene was pumped continuously at the feed rate of 10 ml/hr conversion levels were 20 % for aldehydes (n/i 2.1) after attaining steady state. The reaction was continued for 72 hr without loss of activity. Reaction was arrested by discontinuing the liquid feed and water was pumped for 1 hr. thereafter reactant feed was resumed. Initially there was no conversion, which was steadily resumed over the period of 10 hr. This observation was attributed to formation of water film on the catalyst surface, which physically retards contact of decene with catalyst surface. Moreover water does not wash out complex catalyst, which provides conclusive proof that reaction occurs in the solid state.

The technique of solid catalyst formulation is established in present invention according to which solid catalyst can be formulated and applied for catalyzing reactions in diversity of solvents. The catalytic formulation referred herein was applied to variety of reactions according to yet another embodiment. An exemplary reaction class for which catalytic formulation was employable is described in subsequent sections. Reaction classes that are described here are only exemplary and limited by scope of catalytically active entity as said earlier. Variety of reaction classes described herein are intended to outline the scope of catalytic formulation that is under consideration wherein emphasis given on catalyst separation, stability and convenience of operation when applied to manufacture of plurality of organic compounds. Classes of reaction described herein are hydroformylation, hydrogenation, carbonylation, carbon-carbon bond formation by Heck and Suziki type reactions, isomarization, epoxidation, Wacker oxidation, Michel addition and Knovengel condensation.

Metal catalyzed addition of carbon monoxide and hydrogen to olefin provides access to aldehydes and in certain cases followed by hydrogenation. Wide diversity of olefins can be hydroformylated to corresponding aldehydes various olefins were hydroformylated with analogue of classical Wilkinson's catalyst. Various olefins such as hexene, styrene, cyclohexene and vinylacetate were hydroformylated. Similarly hexene was hydroformylated with rhodium modified with different ligands to control selectivity. Accordingly other metals active for hydroformylation were also tested such as cobalt and platinum.

Palladium phosphine complexes catalyze Carbonylation of halides, alcohols and olefins in homogeneous reaction systems. Analogues of such palladium complexes were formulated and tested for styrene, styryl alcohol, phenyl acetylene and bromobenzene.

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Reasonable activities were obtained in each case and catalyst can be reused.

Various phosphine amines, phosphite complexes of Zr, Hf, V, Cr, Mo, W, Mn, Re, Fe, Ru, Os, Co, Rh, Ir, Ni, Pd, Pt, are useful for hydrogenation of variety of functional groups. Hydrogenation of olefins, carbonyl, and nitroaromatics were tested with the catalyst formulated with this invention.

The extension of substituted alkenes by direct carbon - carbon bond formation at vinylic carbon center is useful reaction for manufacture of variety of organic synthesis. Such synthetic procedure is difficult to achieve by conventional organic synthesis. The palladium complexes (Pd0) have proved to be effective in this sense. Various palladium complexes including phosphine, metalated phosphine and phosphites are useful catalysts other metal such, as nickel and platinum are also useful in this respect. Olefination of aryl bromides was demonstrated with catalytic formulation of the invention.

Palladium catalyzed cross-coupling reactions of aryl or vinyl boronic acids with aryl halides are well known in the art. Such coupling reactions are carried out in polar as well as non-polar media. Palladium phosphine complexes are useful in this respect. Varieties of biaryl compounds are accessible through this reaction. The catalytic formulation of this invention is also suitable for this class of reactions.

Double bond isomarization is useful reaction in converting olefins to isomarized olefins. Various transition metal complexes catalyze this type of reaction. Metal complexes useful in this respect are platinum, palladium, rhodium and cobalt. The catalytic formulation of present invention is also useful in catalyzing this reaction

Present catalytic formulation is also suitable for oxidation of olefins to epoxides and acids. For example molybdate ion when hetereogenized as quaternary ammonium ion pair can catalyze epoxidation of olefins. Various pthalocyanines are also useful in this respect.

Nucleophilic addition of mesomeric anion to activated olefins such as $\alpha\beta$ unsaturated olefins is known as Michel reaction. Compounds containing electron-withdrawing groups having relatively acidic protons are suitable compounds to form mesomeric anions such compounds are for example R-CH₂-Z wherein Z is electron-withdrawing group such as CN, COOR, NO₂, CHO etc. and R may be alky aryl or Z as defined earlier. In presence of strong base these compounds for anion R-CH ^(·)-Z which adds to α - β - unsaturated olefins at β position. The activated olefins may be represented as C=C-Z where in carbon attached to Z is α and adjacent carbon is β .

Generally catalyst employed to form said mesomeric ions are strong bases such as

H, OH, MeO, etc. Anion fragments as such are difficult to solidify therefore counter cation selected for such is quaternary ammonium compounds, which are functionalized with anionic functional groups, and ion pair as a whole is precipitated on solid support. In cases where quaternary ammonium compounds exist as alcoxy ion pair, solidified quaternary compound formulation is successively washed with solution of alcoxy anion prior to use. Condensation of diethyl malonate with ethyl acrylate, diethyl maleate, acrylonitrile, are demonstrated in examples appended hereinafter.

Condensation of aldehydes or ketones, usually not containing an α hydrogen with compounds of the form R-CH₂-Z to form olefins is called as *Knovengel reaction* (Jones, *Org. react.* 1967, 15, 204-599) wherein Z may be CHO, COR, COOH, COOR, CN, NO₂. The catalysts generally employable for this reaction are basic amines, hydroxyl anion or alcoxy anion. Anion fragments as such are difficult to solidify therefore counter cation selected for such is quaternary ammonium compound, which is functionalized with anionic functional groups, and ion pair as a whole is precipitated on solid support. In cases where quaternary ammonium compounds exist as alkoxy ion pair, solidified quaternary compound formulation is successively washed with solution of alcoxy anion prior to use. Condensation of butyraldehyde to 2-ethylhexenal, benzaldehyde and acetone to dibenzyledene acetone, benzaldehyde and acetonitrile to cinnamonitrile are demonstrated in examples appended hereinafter

It would be evident from these descriptions that wide diversity of soluble catalysts can be formulated by appropriately forming catalytic entities that are anionically charged. These entities are structurally analogues to the soluble catalytic entities. The catalysts that are employable in this context are metal complexes, quaternary ammonium compounds wherein complimentary anion is catalyst (complimentary anion can be metal complex, organometallic anion or inorganic anion).

The present invention was conceived without limiting the said solid catalytic formulation to one particular class of complexes or reaction catalyzed by them following certain mechanism, according to some particular theory. It is perceived that solid support having high surface area provides a mechanical strength and a surface upon which insoluble catalytic material is physically implanted. The insoluble material is generated from interaction of otherwise soluble complex catalyst containing two or more anionic functional groups and calcium, strontium and barium salt solutions. This material is formed on the surface of the solid support as a vehicle. The composite solid assembly resulting therefrom can be suitably used as solid catalyst.

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Moreover, recycling and regeneration of applicants' preferred catalytic formulations is readily accomplished using known methods and procedures, for example when acceptable conversion level has occurred in a given batch run whether it is determined by elapsed time or monitored by consumption of substrate or some other parameter. The vessel need only be brought to ambient temperatures and vented off residual pressure if any. The reaction mixture thereafter may simply be separated from catalyst by simple decanting. The catalytic formulation is filtered and possibly washed with suitable liquid for later reuse or simply recharged with feedstock as needed and a subsequent reaction begun.

As catalyst lifetimes are better understood through working with a particular catalyst formulation in repeated recycling in either laboratory or in commercial settings it may be further desirable to regenerate catalyst time to time either by washing with suitable liquid or by specific chemical treatment. Continuos reaction processes are also practicable for applicants preferred catalysts in view of their insolubility and resistance to leaching or other disintegration. Such processes can be designed and implemented using common and known procedures in the art.

For the purpose of further promoting a better understanding of the catalysts and processes of the present invention, reference will now be made in the examples below to specific instances of their preparation and use. These examples are exemplary only and no limitation of the scope or breadth of applicants' invention is intended thereby. Various modifications and variations of this invention will be obvious to a worker skilled in the art and it is to be understood that such modifications and variations are to be included within the purview of this application and the spirit and the scope of the appended claims

Examples

Experiment 1

(Verification of the hypothesis)

This comparative example illustrates the validation of the hypothesis that anions having two or more negative charges when interacted with group IIA metal cations except Mg⁺² invariably result in to a precipitate which is practically insoluble in organic solvents (including nonpolar, polar (protic and aprotic) and sparingly soluble in aqueous solvent in certain cases). This hypothesis was verified as follows. Solutions of different anions were interacted with group IIA metal ions. 0.1 molar solutions of anionic component (solution A) and 0.1 molar solutions of group IIA cation (solution B) component were prepared. 10 ml of solution B was mixed with 50 ml of solution A in boiling tubes solutions were

thoroughly mixed on shaker for 10 hr. resulting suspensions were centrifuged and precipitate was removed by decanting supernatant liquid. Residual precipitate was diluted with distilled water followed by centrifugation and decantation was repeated thrice. To this precipitate 10 ml methanol was added and centrifugation and decantation procedure was repeated resulting wet precipitates were vaccume dried at 50 °C. Mixtures where precipitate was not observed were discarded. Dried precipitate barium and strontium were found insoluble in water, methanol, ethanol, propanol, butanol, acetic acid, benzene xylene, petroleum ether, ethyl acetate, acetone methyl ethyl ketone, acetonitrile, dimethylformamide, chloroform, tetrahydrofuran. Where as some salts of calcium were found sparingly soluble.

Results are summarized in following table.

Solution of group IIA metal cation (0.1molar)	Solution of anion (0.1 molar)	observation
Magnesium chloride	Sodium nitrate	No precipitate
Calcium chloride	Sodium nitrate	No precipitate
Strontium chloride	Sodium nitrate	No precipitate
Barium chloride	Sodium nitrate	No precipitate
Magnesium chloride	Sodium propionate	No precipitate
Calcium chloride	Sodium propionate	No precipitate
Strontium chloride	Sodium propionate	No precipitate
Barium chloride	Sodium propionate	No precipitate
Magnesium chloride	p-toluene sulfonate sodium	No precipitate
Calcium chloride	p-toluene sulfonate sodium	No precipitate
Strontium chloride	p-toluene sulfonate sodium	No precipitate
Barium chloride	p-tolune sulfonate sodium	No precipitate
Magnesium chloride	m- benzene disulfonate disodium	No precipitate
Calcium chloride	m- benzene disulfonate disodium	White precipitate
Strontium chloride	m- benzene disulfonate disodium	White precipitate
Barium chloride	m- benzene disulfonate disodium	White precipitate
Magnesium chloride	di-sodium oxalate	No precipitate
Calcium chloride	di-sodium oxalate	White precipitate
Strontium chloride	di-sodium oxalate White precipitate	
Barium chloride	di-sodium oxalate White precipitate	
Magnesium chloride	Sodium sulfate	No precipitate

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Calcium chloride	Sodium sulfate	White precipitate
Strontium chloride	Sodium sulfate	White precipitate
Barium chloride	Sodium sulfate	White precipitate
Magnesium chloride	di-sodium phenyl phosphonate	No precipitate
Calcium chloride	di-sodium phenyl phosphonate	White precipitate
Strontium chloride	di-sodium phenyl phosphonate	White precipitate
Barium chloride	di-sodium phenyl phosphonate	White precipitate
Magnesium chloride	Na ₂ HPO ₄	No precipitate
Calcium chloride	Na₂HPO₄	White precipitate
Strontium chloride	Na ₂ HPO ₄	White precipitate
Barium chloride	Na ₂ HPO ₄	White precipitate
Magnesium chloride	di-sodium phthalate	No precipitate
Calcium chloride	di-sodium phthalate	White precipitate
Strontium chloride	di-sodium phthalate	White precipitate
Barium chloride	di-sodium phthalate	White precipitate
Magnesium chloride	Ammonium molybdate	No precipitate
Calcium chloride	Ammonium molybdate	White precipitate
Strontium chloride	Ammonium molybdate	White precipitate
Barium chloride	Ammonium molybdate	White precipitate
Magnesium chloride	Sodium carboxy methyl cellulose	No precipitate
Calcium chloride	Sodium carboxy methyl cellulose	White precipitate
Strontium chloride	Sodium carboxy methyl cellulose	White precipitate
Barium chloride	Sodium carboxy methyl cellulose	White precipitate
Magnesium chloride	Sodium polyvinyl sulfonate	No precipitate
Calcium chloride	Sodium polyvinyl sulfonate	White precipitate
Strontium chloride	Sodium polyvinyl sulfonate	White precipitate
Barium chloride	Sodium polyvinyl sulfonate	White precipitate

In addition to these, EDTA acetyl acetonate, hydride compounds of calcium were interacted with sodium sulfate, sodium phosphate to yield insoluble precipitate.

Experiment 2

Synthesis of high purity oleum

2 lit. Three-necked flask was attached with distillation condenser, addition funnel and to another end collection vessel with bottom drain valve. Distillation condenser was also provided with pressure relief non-return valve. In flask magnetic bar was placed and

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500 g. P₂O₅ was charged. 45 ml conc. H₂SO₄ was placed in collection vessel. 400 ml conc. H₂SO₄ was placed in addition funnel. Addition was started with simultaneous magnetic agitation over the period of 2 hours. Temperature of flask was raised slowly until slow distillation of sulfur trioxide was started. Sulfur trioxide was collected in conc. H₂SO₄ in collection vessel after total volume of liquid in collection vessel reached to 148 ml, heating was stopped and assembly was dismantled.

Experiment 3

Synthesis of Triphenyl phosphine trisulfonate

Triphenyl phosphine trisulfonate was synthesized by following procedure. Triphenylphosphine 50 g. was placed in sulfonation reaction followed by vaccume argon degassing and blanketed with argon. Sulfonation reactor was cooled to 5 °C and 200 g sulfuric acid was charged in the sulfonation reactor without allowing temperature of reactor to cross 10 °C. Addition of sulfuric acid was carried out with constant stirring with mechanical stirrer over a period of 2 hours. Reaction mixture assumed pale yellow color. To this reactor 280 g of 65 % oleum prepared as per previous experiment was charged over a period of 60 min. temperature of the sulfonation reactor was raised to 22 °C and reaction was continued for 76 hours. There after temperature of the reaction was lowered to 0°C and 50 ml distilled and degassed water was introduced in the sulfonation reactor without allowing temperature to rise beyond 5 °C over a period of three to four hours. This solution was further diluted with 500 ml water. The diluted solution was transferred to 3-lit jacketed vessel and chilled to 5 °C and consequently neutralized with 50 % w/w NaOH in water, which was previously degassed. At neutralization point solution assumed distinct yellow color at this instance NaOH addition was discontinued and pH was lowered to 6 by addition of con sulfuric acid. During neutralization formed sodium sulfate partially precipitates which was removed by filtration and resulting solution was concentrated under vacuum to 300 ml. formed sodium sulfate was removed by filtration. Mother liquor containing TPPTS was further diluted with 2000 ml degassed methanol and refluxed for two hours during which most of the sodium sulfate precipitated, supernatant extract of TPPTS in methanol was removed by filtration TPPTS extract in methanol was evaporated to dry ness and white colored solid was obtained (purity above 95% by P31NMR). This solid was dissolved in minimum amount of water and reprecipitated with degassed ethanol to obtain TPPTS with purity > 99 %.

Experiment 4

Synthesis of disodium P-phenyl-3, 3'-phosphine dialy bis (benzene sulfonate)

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Orthoboric acid (48 g) was dissolved in concentrated sulfuric acid 98% (200 ml) to this was added 65% oleum 200 ml. the temperature of the solution was raised to 60°C and excess sulfur trioxide was removed in high vaccume by providing a gas trap attachment containing calcium oxide (trap was chilled to -10 °C) solution of orthoboric acid and sulfur trioxide was cooled to 5°C and 30 g triphenyl phosphine was added under argon blanket. Resulting mixture was agitated by mechanical stirrer and temperature of the reactor was raised to 58°C and reaction was continued for 90 hours. The temperature was reduced to 0°C and hydrolyzed with 500 ml degassed water. This solution was neutralized with 50% w/w sodium hydroxide in water until neutralization and formed precipitate was removed by filtration and mother liquor was concentrated to 300ml and diluted with 1000 ml methanol and refluxed for 2 hours. Resulting precipitate was removed by filtration. The extract in methanol was evaporated to obtain a solid which was suspended in 1000 ml methanol and to this 50 g microcrystalline cellulose avicel was added followed by 20 ml conc. H₂SO₄ and refluxed for 6 hours under argon blanket. Solution was cooled and filtered to remove avicel. To this 50 g. Avicel™ was again added and refluxed for another 6 hours suspension was filtered and methanolic extract was neutralized with 50 % NaOH w/w and filtered. Solution was evaporated to obtain white compound correct elemental analysis.

Experiment 5

Synthesis of trisodium 3,3', 3''-phosphine trial tris (4 methyl benzene sulfonate)

Orthoboric acid (g., 6.6 mmol) was dissolved in concentrated sulfuric acid (96%, 3.75 ml) to this (2-methylphenyl) phosphine (0.50 g., 1.4 mmol) was dissolved in reaction mixture. Oleum (6.75 ml, 65 % w/w) was added drop wise while temperature of reaction mixture was 0 °C. After stirring for 72 hours at 25 °C, the temperature was lowered to 0 °C and the mixture was hydrolyzed by addition of 5 ml degassed water. This solution was neutralized with 50 % w/w sodium hydroxide in water until neutralization and formed precipitate was removed by filtration and mother liquor was concentrated to 3 ml and diluted with 10 ml methanol and refluxed for 2 hours. Resulting precipitate was removed by filtration. The extract in methanol was evaporated to obtain a solid which was suspended in 10 ml methanol and to this 0.5 g microcrystalline cellulose avicel was added followed by 0.5 ml conc. H₂SO₄ and refluxed for 6 hours under argon blanket. Solution was cooled and filtered to remove avicel. To this 0.5 g. aviel was again added and refluxed for another 6 hours suspension was filtered and methanolic extract was neutralized with 50

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% NaOH w/w and filtered. Solution was evaporated to obtain white compound correct elemental analysis.

Experiment 6

Synthesis of sodium salt of sulfonated tribenzyl phosphine

Sulfonation of tribenzyl phosphine was carried out anoaloguous to triphenyl phosphine except exact degree of sulfonation was not established reaction mixture containing di and tri sulfonated phosphine was used for further experiments.

Experiment 7

Sulfonation of 1-3 bis -diphenyl phosphino propane

4.95 g. (12 mmol) of diphenylphosphino propane was dissolved in a solution of 4 g. (64.7 mmol) orthoboric acid in 37.5 ml (98%) reaction mixture was cooled to 0 °C, to this 65% oleum 67.5 ml was added drop wise over a period of 2 hours. After addition reaction mixture was brought to 25 °C and stirred for 48 hours. After this reaction mixture was brought to 0 °C and hydrolyzed with 50 ml degassed water. This solution was neutralized with 50 % w/w sodium hydroxide in water until pH 7 and formed precipitate was removed by filtration and mother liquor was concentrated to 30 ml and diluted with 100 ml methanol and refluxed for 2 hours. Resulting precipitate was removed by filtration. The extract in methanol was evaporated to obtain a solid which was suspended in 100 ml methanol and to this 5 g microcrystalline cellulose avicel was added followed by 1 ml conc H₂SO₄ and refluxed for 6 hours under argon blancket. Solution was cooled and filtured to remove avicel. To this 5 g. avicel was again added and refluxed for another 6 hours suspension was filtered and methanolic extract was neutralized with 50 % NaOH w/w and filtered. Solution was evaporated to obtain a white compound

Experiment 8

Sulfonation of 1-2 bis-diphenyl phosphino ethane

Preparation was carried out in analogous manner as explained in previous experiment

Experiment 9

2,2'-bis(diphenylphosphinomethyl)-1,1'biphenyl (Bisbi) synthesis and sulfonation

Equip a three litre three-necked flask with a sealed mechanical stirrer, a reflux condenser and a thermometer. Dissolve 89 g. (0.5 mol) of phenanthrene in one litre glacial acetic acid in the flask and warm to 85°C on a water bath. Introduce 345 ml of 30% hydrogen peroxide solution (4 mol) during 40 min. temperature falls to about 80°C continue for 6 hr. remove acetic acid and water under reduced pressure to obtain brown color solid digest this residue in 2N sodium hydroxide solution and add 4 g of powdered

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charcoal and warm the mixture to 75°C and filter. Filtrate was acidified to pH 2 with the conc. HCl white precipitate was filtered and dried at 50°C MP 109°C, 83 g 69 % material obtained is of sufficient purity for further synthesis.

Equip a three litre three-necked flask with a sealed mechanical stirrer, a reflux condenser and a thermometer. Flask was cooled to 0 °C in ice salt bath. Reaction vessel was charged with 24.2g (0.1 mol) diphenic acid and 15.12 g (0.4 mol) sodium borohydride to this solid powder 200 ml dry tetrahydro furan was added in such a way that there is minimum effervescence. After 1 hour suspension becomes uniform and to this (0.2 mol H₂SO₄ in 100 ml tetrahydrofuran was added over a period of 2 hours while maintaining temperature at 0 °C. after addition was over mixture was allowed to stirr for 24 hours at room temperature. To this white suspension 100 ml 30 % NaOH was added and refluxed for 4 hours and liquid was braught to room temperature and extracted with chloroform to yield white solide. Which was used further without purification.

Diol intermediate (0.08 mol) from above said preparation was dissolved in chloroform and transferred to two necked flask attached with condenser and guard tube, pressure equalizing addition vessel. One drop of pyridine was added to flask and (0.2 mol) thionyl chloride was dissolved in 25 ml chloroform and charged in addition vessel. Thionyl chloride was added to round bottom flask at room temperature. During addition considerable amount of sulfur dioxide and hydrogen chloride escaped from guard tube. The temperature of the flask was raised until chloroform started refluxing. After 5 hours reaction was quenched by addition of water. Chloroform was exteracted with bicarbonate solution followed by water and dried by passing through bed of sodium sulfate. Chloroform was evaporated under vaccume at 50 °C to yild yellow colored oil (irritant and inflammatory to skin), which was distilled, in high vaccume to yield pale yellow colored oil.

(Following procedure was adopted from US patent 4,879,416).

To a 500ml flask equipped with a mechanical stirrer, thermowell, addition funnel and condencer was added triphenyl phosphine 16.77 g, 0.064 mols, tetrahydrofuran 64 ml and lithium wire 0.88 g, 0.128 atoms. The flask was cooled to 15 °C reaction mixture was stirred overnight to yield red colored solution with complete dissolution of lithium. The flask was cooled further to 5 °C and tertiary butyl chloride 5.92 g 0.064 mols was added and temperature was raised to 50 °C and maintained for 2 hours. Reaction mixture was cooled and to this 7.5 g of above said dichloride was slowly added. Temperature of the reaction mixture was raised such that it gently boils. Reaction was quenched by addition of

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5 ml methanol. The reaction mixture was evaporated to yield sticky mass, which was dissolved in sufficient diethyl ether and washed with water. Evaporation of diethyl ether yields pale yellow colored sticky mass, which was recrystallized from THF/Methanol to yild fine crystals of white material.

This material was sulfonated according to method described for diphenyl phosphino propane. To produce white coloured compound, which was soluble in water.

Experiment 10

Sulfonation of (R) BINAP (2, 2'-bisdiphenylphosphino - 1,1' binapthyl)

Procedure of sulfonation was adopted from US patent 5756838. 0.5 g. of (R) BINAP was dissolved in 1.75 ml of concentrated sulfuric acid at 10 °C under argon. Afterwards, 7.5 ml of oleum 40 % w/w was added dropwise over 2-3 hours the resulting mixture was stirred at 10 °C for 76 hours. After stirring this mixture was slowly poured over 100 g ice followed by dropwise addition of 50 % w/w NaOH untill solution was neutralized to pH 7. The resulting solution was concentrated under vaccume to 30 ml. to this 100 ml methanol was added in order to precipitate sodium sulfate. Methanolic extracted was evaporated under vaccume to obtain solid, which was dissolved in methanol and filtered. Methanol was evaporated to obtain white solid.

Similarly s BINAP was sulfonated.

Experiment 11

Sulfonation of (S, S chiraphos) (S) (S) 2,3 bisdiphenylphosphino butane.

Procedure of sulfonation was adapted from Alario et al, J. Chem. Soc., Chem. Commun., 1986,202

Experiment 12

Sulfonation of R prophos 1, 2(S) bisdiphenylphosphino propane

Procedure of sulfonation was adapted from Amrani et al Organometallics 1989, 8, 542

Experiment 13

Sulfonation of R, R 2-5, bis diphenylphosphino penatne

Procedure of sulfonation was adapted from Amrani et al Organometallics 1989, 8, 542 Sulfonation of 2-pyridyl phosphine

Experiment 14

Synthesis of sodium salt of sulphonate of triphenylamine.

2 g of Triphenyl amine was charged into a reactor, and 20 cc of concentrated sulfuric acid was added to it. This mixture was stirred until the amine dissolved. 20 cc of oleum 65% was added to this mixture under rapid stirring, and the reactor was cooled to

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about 20°C. After the addition of oleum, the reactants and contents were heated to 50°C and maintained at this temperature for 48 hours. The reactor and its contents were cooled, and distilled water (10cc) was added to the reaction mixture to quench the oleum. 50% NaOH solution was added to this solution, under cooling (10 °C) until the sulfuric acid solution was neutralized. The solution was concentrated and then methanol was added to extract the water-soluble ligand from the sodium sulfate powder. The methanol was evaporated to yield the water-soluble sodium salt of triphenyl amino sulfonic acid [1.6g]. The product consists of mixtures of the bis and grater than 95 % tris sulfonation products. These can be used as such in synthesis of metal complexes for catalysis.

Experiment 15

Trisodium salt of tribenzylamine trisulfonate

2 g of Tribenzyl amine was charged into a reactor, and 20 cc of concentrated sulfuric acid was added to it. This mixture was stirred until the amine dissolved. 20 cc of oleum 65% was added to this mixture under rapid stirring, and the reactor was cooled to about 20° C. After the addition of oleum, the reactants and contents were heated to 50°C and maintained at this temperature for 48 hours. The reactor and its contents were cooled, and distilled water (10cc) was added to the reaction mixture to quench the oleum. 50% NaOH solution was added to this solution, under cooling (10 ° C) until the sulfuric acid solution was neutralized. The solution was concentrated and then methanol was added to extract the water-soluble ligand from the sodium sulfate powder. The methanol was evaporated to yield the water-soluble sodium salts of Tribenzyl amino sulfonic acid [1.7 g] degree of sulfonation was established by H1 NMR and elemental analysis.

Experiment 16

Synthesis of sodium salt of sulphonate of 2,2'bipyridine.

2 g of 2,2'Bipyridine was charged into a reactor, and 20 cc of concentrated sulfuric acid was added to it. This mixture was stirred until the amine dissolved. 20 cc of oleum 65% was added to this mixture under rapid stirring, and the reactor was cooled to about 20°C. After the addition of oleum, the reactants and contents were heated to 50°C and maintained at this temperature for 48 hours. The reactor and its contents were cooled, and distilled water (10cc) was added to the reaction mixture to quench the oleum. 50% NaOH solution was added to this solution, under cooling (10 °C) until the sulfuric acid solution was neutralized. The solution was concentrated and then methanol was added to extract the water-soluble ligand from the sodium sulfate powder. The methanol was evaporated to yield the water-soluble sodium salt of 2,2'bipyridine di sulfonic acid. [1.2 g] The product

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consists of mixtures of the bis sulfonation products as indicated by elemental analysis and 1h NMR. These can be used as such in synthesis of metal complexes for catalysis.

Experiment 17

Sulfonation of 2 phenyl pyridine

2 g of 2,phenylpyridine was charged into a reactor, and 20 cc of concentrated sulfuric acid was added to it. This mixture was stirred until the amine dissolved. 20 cc of oleum 65% was added to this mixture under rapid stirring, and the reactor was cooled to about 20°C. After the addition of oleum, the reactants and contents were heated to 50°C and maintained at this temperature for 48 hours. The reactor and its contents were cooled, and distilled water (10cc) was added to the reaction mixture to quench the oleum. 50% NaOH solution was added to this solution, under cooling (10 °C) until the sulfuric acid solution was neutralized. The solution was concentrated and then methanol was added to extract the water-soluble ligand from the sodium sulfate powder. The methanol was evaporated to yield the water-soluble sodium salt of 2-phenylpyridine sulfonic acid. [1.2 g] The product consists of mixtures of the bis sulfonation products as indicated by elemental analysis. These can be used as such in synthesis of metal complexes for catalysis.

Experiment 18

Synthesis of 2-3 bisdiphenylphosphino, succinic acid sodium salt

To a reaction system comparising a solution of dimethyl maleate (50 g.) in chloroform (100 ml) was added a solution of bromine (15 ml) in chloroform 100 ml over a period of 2 hours. The reaction mixture was stirred for 2 hours at the end of reaction mixture was washed twice with 100 ml saturated sodium thiosulphate and then twice with 100 ml water. Organic part was passed through 5 g. sodium sulphate and sub sequently treated with activated charcoal. Chloroform was stripped off to yield 60 g oil.

Subsequent reaction was set up with 250 ml. three necked glass vessel equipped with addition funnel magnetic stirrer and rubber septum. Assembly was flushed with argon. To this vessel finely cut lithium ribbon (500 mg.) was added and assembly was evaccuated and refilled with argon. To this assebly 50 ml tetra hydrofuran was added with gas tight syringe maintaining argon blanket 8.3 ml chlorodiphenyl phosphine was placed in addition funnel set up was evacuated and refilled with argon contents of addition funnel were dropped in the lithium suspension, during lithium dissolution solution started assuming red color and reaction mixture was stirred for 4 hours after complete dissolution of lithium.

To another 250 ml vessel equipped with reflux condenser and rubber septum 30 ml dry tetrahydrofuran was placed by syringe and assembly was evaccuated and refilled with

argon. To this 4.52 g of brominated diethyl maleate was transferred by syringe followed by 30 ml of lithium phosphied soultion (red colored). Contents of the reaction mixture were maintained at 80 °C for 12 hours. To this reaction mixture 1 ml methanol was added and tetrahydrofuran was removed under vacuum. Syrupy orange coloured liquid was washed twice with 25 ml ether. 1 g of this syrupy orange product was transferred to three necked round bottomed flask attached with reflux condencer the set up was thoroughly flushed with argon and 20 ml 2 % sodium hydroxide were refluxed the reaction mixture was cooled to 5°C and precipitated white material of diphosphine was recovered by filtration yield 1 g.

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Experiment 19

Quaternization of tribenzyl amine tri sulfonate with benzyl chloride

To a mixture of (0.1 mol) tribenzylamine trisulfonate and benzyl chloride (0.2 mol) was added 50 ml water and 50 ml dimethyl formamide. Solution was stirred at 70 °C for 76 hours and reaction was monitored by disappearance of benzyl chloride. Reaction mixture was evaporated under vacuum to yield a solid mass, which was dissolved in minimal amount of water, and aqueous solution was washed with diethyl ether. Aqueous extract was dried under vacuum and solid was stored in dry condition.

Experiment 20

Synthesis of quaternary ammonium hydroxide

17 g. (0.1 mol) of silver nitrate was dissolved in 170 ml of distilled water and warmed to 85°C and 3.9g (0.097 mol) sodium hydroxide was added to it. Mixture was agitated vigorously until coagulation of precipitation is complete. Precipitate was recovered by centrifugation and suspended in 100 ml water to which was added (0.09 mol) of above quaternaryammonium compound. Reaction mixture was stirred for 3 hours under nitrogen and filtured. Liquid was evaporated under vacuum at room temperature to obtain a solid.

Experiment 21

Quaternization of triphenyl amine with benzyl chloride

To a mixture of (0.1 mol) triphenylamine trisulfonate and benzyl chloride (0.2 mol) was added 50 ml water and 50 ml dimethyl formamide. Solution was stirred at 70°C for 76 hours and reaction was monitored by disappearance of benzyl chloride. Reaction mixture was evaporated under vacuum to yield a solid mass, which was dissolved in minimal amount of water, and aqueous solution was washed with diethyl ether. Aqueous extract was dried under vacuum and solid was stored in dry condition.

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Experiment 22

Formation of quaternary ammonium hydroxide of quaternary ammonium salt of n benzyl triphenyl amine.

Experiment 23

5 Synthesis of hydridocarbonyl tris (trisodiumtriphenylphosphine trisulfonate) rhodium (I)
The procedure was adopted from US patent 4, 994,427 dated Feb. 19, 1991 to Davis et al.
500 mg. Acetyl acetonate dicarbonyl rhodium (I) was added to vigorously stirred 10 ml
deaerated solution of 4 g. of sodium triphenylphosphine trisulfonate in water. After
dissolution was complete stirring was continued for six hours under atmosphere of 1;1
H₂/CO. solution was then centrifugend to remove precipitated rhodium. To this solution 80
ml absolute ethanol saturated with 1:1 H₂/CO were added to precipitate desired complex.
Precipitate was recovered and vaccume dried.

Experiment 24

Dichloro bis (tris triphenylphosphine sulfonato trisodium) palladium (II)

This procedure was adapted from Jiang et al J. Mol. Catal. A: Chemical 130 (1998) 79-84, 100 mg PdCl₂ and 2 ml 2 M HCl were added to a schlenk flask and the mixture was stirred at 50°C until PdCl₂ was dissolved completely. After the flask was cooled to room temperature and flushed with argon, 0.80 g. TPPTS was added in to the flask under stirring. The color of the solution changed from dark red to yellow immediately. After 10 min stirring, 15-ml ethanol was added, alight yellow powder precipitated and mixture was stirred for 30 min. The filtered precipitate was washed three times with 30-ml. warm 95 % ethanol and dried in vacuum.

Experiment 25

Synthesis of trans-PtCl₂ (TPPTS)₂

The platinum complex PtCl₂ (NCPh)₂ 235 mg (0.5 mmol) was dissolved in 10 ml toluene to this solution was added to aqueous solution of TPPTS (568 mg 1 mmol) in 10 ml water to this mixture isopropanol 3 ml was added and reaction mixture was stirred at 50 °C for 10 h complex was recovered from aqueous phase by evaporation 620 mg of PtCl₂ (TPPTS)₂. 6 H₂O.

Experiment 26

Synthesis of NiCl₂/ TPPTS

Nickel chloridehexahydrate (0.05 mols) was reacted with tppts (0.12 mols) in water sufficient to dissolve and formed complex was precipitated by ethanol

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Experiment 27

Syntheis of IrCl (COD) / TPPTS

IrCl (COD) (0.01 mol) was dissolved in minimum amount of tolune and exchanged with). 04 mols of tppts dissolved in minimum amound of water. Tolune layer was removed and aqueous layer was dried.

Experiment 28

Synthesis of [Ru (Cl) (μ -Cl) (TPPTS)₂]

The method was adopted from M. Hernandez et al, J. Mol. Catal. A: Chemical 116 (1997) 117-130. RuCl₂ (PPh₃)₃ 5.8 g. 6 mmol was dissolved in 150 ml of tetrahydrofuran and heated to 60 °C. A 30 ml water solution of TPPTS (6.3g 10.1 mol was added drop wise under vigorous stirring. The biphasic medium was stirred further for 30 min at 60 °C. After cooling to room temperature, 140 ml of orange organic layer was removed. The resulting solution was filtered out. Then the deep red aqueous phase was evaporated to dryness and further dried in vacuum.

Experiment 29

Synthesis of [Ru(H)(Cl)(TPPTS)₃]

The method was adopted from M. Hernandez et al, J. Mol. Catal. A: Chemical 116 (1997) 117-130. This complex was prepared from [Ru(H)(Cl)(PPh₃)₃]. PhCH₃ 3 g. 3.3 mmol dissolved in 120 ml tetra hydrofuran; TPPTS 5 g. (8 mmol); H₂O 30 ml. a bright purple coloured solid was recovered from aqueous layer.

Experiment 30

Synthesis of [Ru(H)₂(TPPTS)₄]

0.1 g. (0.38 mmol) of RuCl₃. 3H₂O and 1.07 g TPPTS 1.72 mmol were dissolved in 10 ml of distilled water. The deep brown coloured solution was stirred at room temperature while passing stream of hydrogen. After 10 min 0.17 g. (~4.5 mmol) of NaBH₄were added. Solution turned instantaniously brown yellow with vigorous effervescence. The mixture was heated to 50 °C for 10 min after cooling and evaporation to dryness solid was obtained

Experiment 31

Synthesis of Ru/Binapts complex

Ruthenium binap 4 SO₃Na catalyst was prepared by reacting (0.01 g) of [Ru(benzene)Cl₂]₂ with two equivalents of (0.05 g) R- binap 4 SO₃Na in a 1:8 benzene ethanole mixture 4.5 ml to yield [Ru(benzene)Cl] R- binap 4 SO₃Na. Resulting solution was vaccume dried

Experiment 32

Synthesis of Rh[†]/ chiraphos tetra sulfonate complex

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Rh⁺/ chiraphos tetra sulfonate catalyst was prepared by reacting [Rh (COD) Cl]₂, with two mole equivalents of sulfonated ligand in water at room temperature in presence of excess sodium perchlorate to form cationic complex

Exaperiment 33

Synthesis of palladium acetate sulfonated bypyridyl complex

Synthesis procedure was adopted from brink et al Chem. Commun, 1998, 2359-2360. Pd(OAc)₂ 0.1 mmol and sulfonated bypyridyl 0.1 mmol were stirred overnight with 42.5 g of water to afford a clear orange colored solution which was evaporated to dryness.

Experiment 34

Tetrasodium salt of Cobalt (II) 4,4', 4", 4", - Tetrasulfophthalocynine (procedure is adopted from Inorg. Chem. Vol 4, No. 4 April 1965, 469-471)

The monosodium salt of 4-sulfopthalic acid (4.32 g., 0.0162 mol.), ammonium chloride (0.47 g., 0.009 mol.), urea (5.8 g., 0.097 mol.) ammonium molybdate (0.068g., 0.00006 mol), and cobalt (II) sulfate 2 H₂O (1.36 g., 0.0048 mol) and 100 ml celite were ground together in nitrobenzene to form a homogeneous paste and diluted to 50 ml with nitrobenzene in round bottomed flask attached with reflux condenser. The reaction mixture was heated to 180 °C. The reaction mixture was heated slowly with overhead stirring while maintaining temperature 180 - 190 °C. The heterogeneous mixture was heated for 6 hours at 180 °C. The crude product was recovered by cooling reaction mixture and removing nitrobenzene. Solid cake was washed with hexane followed by methanol until nitrobenzene was removed. The solid residue was transferred to 110 ml 1 N hydrochloric acid saturated with sodium chloride. The mixture was heated briefly to boiling, cooled to room temperature and filtered. The resulting solution was dissolved in 70 ml of 0.1 N NaOH. The solution was heated to 80 °C and insoluble impurities were immediately separated by filtration. Sodium chloride (27 g. was added to solution and slurry was heated to 80 °C until ammonia evolution ceased. Reaction mixture was cooled to room temperature and filtered. This re-precipitation process was repeated twice and solid was filtered and washed with 80 % ethanol until filtrate was chloride free as tested by silver nitrate solution. This solid was refluxed in 20-ml ethanol for 4 hours to get pure product, which was dried over P₂O₅ yield 65 %correct elemental

Experiment 35

Tetrasodium salt of copper(II) 4, 4', 4", 4", - Tetrasulfophthalocynine

The compound was prepared using similar mole ratios of the reactant except 0.0048 mol

of copper sulfate $5.H_2O$. and purified as described for Tetrasodium salt of Cobalt(II)4,4', 4''', - Tetrasulfophthalocynine

Experiment 36

Tetrasodium salt of Manganese(II) 4, 4', 4", 4", - Tetrasulfophthalocynine

The compound was prepared using similar mole ratios of the reactant except 0.0048 mol of manganese acetate and purified as described for Tetrasodium salt of Cobalt(II)4,4', 4'', 4''', - Tetrasulfophthalocynine

Experiment 37

Tetrasodium salt of iron(III) 4, 4', 4'', 4''', - Tetrasulfophthalocynine oxygen adduct
The compound was prepared using similar mole ratios of the reactant except 0.0048 mol of
Fe (III) chloride and purified as described for Tetrasodium salt of Cobalt (II) 4,4', 4'', 4''',
- Tetrasulfophthalocynine.

Experiment 38

Water soluble cobalt II complex N, N' – ethylenebis (salycyldiamine 5- sodium sulfonate) Synthesis of this comples was performed according to Kevin et al. J. Chem. Soc., Dalton Trans. 1982, 109.

N- phenyl salicyldimine (35 g.) was added to concentrated sulfuric acid 95 cm³ and mixture was heated for two hours, with occasional stirring while keeping temperature in the range of 100 +-5 °C and after cooling solution was slowly poured over ice water to obtain yellow precipitate which was subsequently recrystallized from water to obtain crystalline yellow compound (20 g).

25.5 g of above product was dissolved in 500 ml water and to this solution 8.4 g. unhydrus sodium carbonate was slowly added and stirred until effervescence ceased aniline was steam distilled aqueous solution was vaccume dried to obatin a solid which was purified by precipitation from water and ethanol.

Na₂[Co (SO₃sal)]. 3 H₂O

- 30 The compound CoCl₂. 6H₂O (6 g. 25 mmol) was dissolved in 30 cc ater and added to a solution of disodium salicsyldehyde 5 sulfonic acid (13.2 g 50 mmol) in 20 cc water and mixture was heated for 10 min. After filtration solution was concentrated and cooled to obtain 12 g of crystalline complex.
- N, N' ethylenebis (salycyldiamine 5- sodium sulfonate)

 Ethanol 100cc water 15 cc and ethylene diamine (0.6 g. 10 mmol) were added to Na₂ [Co

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(SO₃sal)]₂. 3 H₂O (5.5 g. 10 mmol and mixture was refluxed under nitrogen atmosphere for 1 hr. dark brown feathery precipitate was recovered

Experiment 39

Preparation of supports for catalyst preparation

All support materials were sourced from commercial suppliers and were used without further size reduction. Specifications of supports are provided with appropriate specifications. Support materials were extracted with hexane, ether methanol and water using assembly described in figure 3.

Surface saturation with group IIA ions

- 10 Each support was divided in to a lot of 25 g and suspended in 500 ml solution of 5 % barium nitrate solution. The suspension was refluxed for 24 hours. Suspension was brought to room temperature and solid were filtered and transferred to extractor described in figure 3 and extracted with 500 ml of water, acetone and petroleum ether (bp 60- 80 °C) solids were vacuum dried and stored for further use.
 - Degassing supports as described above were degassed immediately before use by following procedure. Required amount was transferred to round bottomed flask equipped with two-way valve and evacuated at 0.1 mm Hg and temperature was raised to 150 0 C and kept at this temperature for 1 hour at this temperature while maintaining vacuum. Vacuum inlet was closed and argon was introduced and flask was cooled to room temperature. The procedure was repeated at least thrice and solid was stored under argon for further use.

Following supports were prepared accordingly, silica, gamma alumina, zirconia, titania, keisulghur, bentonite, hyflosupercel, asbestos powder, magnesium hydrotalcite, barium sulfate, charcoal, bone ash.

Example 1 to 84

25 Preparation of catalytic formulation by co-precipitation

The following examples illustrate one of the procedures for the preparation of the catalytic formulation of the invention in accordance with the method of formulation known as co precipitation in bulk liquid.

The general procedure for the preparation of heterogeneous catalytic formulation is described herein as making of a solution of anionically charged catalytic entity, catalytically inert anionic additive (termed as solution A) and solution of group II A metal ions (termed as solution B). A support pretreated as described in earlier is suspended in aqueous or water miscible solvent and resulting suspension is vigorously agitated to this suspension solution A and solution B were added over a prolonged period of time and

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resulting suspension is further agitated for specified time. Suspension was centrifuged and solids were repeatedly washed with water, methanol and diethyl ether followed by drying in vacuum. Dry powder was stored under argon in gas tight vessel and can be used for appropriate reaction depending upon catalytically active entity incorporated in it.

Note1: solution A is prepared by dissolving anionic components including anionic complex and additives to make homogeneous solution in degassed solvents. The resulting solution is also degassed by purging argon.

Note 2: solution B is prepared by dissolving dissolving group IIA metal salts.

Solution was degassed prior to use.

Note 3 addition of A and B is carried out at ambient temperature unless stated.

Example	Solution A	Solution B	Procedure
1	HRhCO(TPPTS)3, 50 mg,	Saturated barium	A suspension of 2 gm Davisil in 10 ml water was
	TPPTS 200 mg.	nitrate in water 2 ml	formed and resulting suspension was vigorously
,	Dissolved in water 2 ml		agitated to this suspension solution A and solution B
	·		were added simultaniously over a 3 hours in 50 μl
			portions resulting suspension is further agitated for 10
			hours to yild pale yellow colored solid powder.
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2	HRhCO(TPPTS) 3, 50 mg,	Saturated strontium	Solution A and solution B were added to a suspension
	TPPTS 200 mg.	chloride in water 2ml	of 2 gm Davisil in 10 ml water and resulting suspension
	Dissolved in water 2 ml	·	is vigorously agitated to this suspension solution A and
	,		solution B were added simultaniously over a 3 hours in
			50 μl portions resulting suspension is further agitated
			for 10 hours to yild pale yellow colored solid powder.
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·			
3	HRhCO(TPPTS)3, 50 mg,	500 mg of calcium	Solution A and solution B were added to a suspension
	TPPTS 200 mg.	chloride in 2 ml	of 5 gm Davisil in 10 ml water and resulting suspension
	Dissolved in water 2 ml	water	is vigorously agitated to this suspension solution A and
			solution B were added simultaniously over a 3 hours in
			50 µl portions resulting suspension is further agitated
			for 10 hours to yild pale yellow colored solid powder.

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4	HRhCO(TPPTS)3 50 mg,	Barium nitrate	Solution A and solution B were added to a suspension
	TPPTS 200 mg.	saturated solution in	of 2 gm γ-alumina in 10 ml water and resulting
	Dissolved in water 2 ml	water	suspension is vigorously agitated to this suspension
		•	solution A and solution B were added simultaniously
			over a 3 hours in 50 µl portions resulting suspension is
		•	further agitated for 10 hours to yild pale yellow colored
			solid powder
5	HRhCO(TPPTS)3, 50 mg,	Strontium chloride	Solution A and solution B were added to a suspension
	TPPTS 200 mg.	saturated solution in	of 2 gm γ-alumina in 10 ml water and resulting
İ	Dissolved in water 2 ml	water	suspension is vigorously agitated to this suspension
			solution A and solution B were added simultaniously
			over a 3 hours in 50 µl portions resulting suspension is
			further agitated for 10 hours to yild pale yellow color
			solid powder
6	HRhCO(TPPTS)3 50 mg,	Calcium chloride 500	Solution A and solution B were added to a suspension
1		mg solution in 2 ml	of 2 gm γ-alumina in 10 ml water and resulting
	TPPTS 200 mg.	water	suspension is vigorously agitated to this suspension
	Dissolved in water 2 ml	,	solution A and solution B were added simultaniously
		•	over a 3 hours in 50 µl portions resulting suspension is
			further agitated for 10 hours to yild pale yellow color
	<u> </u>		solid powder
7	HRhCO (TPPTS)3, 50 mg,	Barium nitrate	Solution A and solution B were added to a suspension
	TPPTS 200 mg.	saturated solution in	of 2 gm bentonite in 10 ml water and resulting
	Dissolved in water 2 ml	water	suspension is vigorously agitated to this suspension
	Dissolved in water 2 in	water	solution A and solution B were added simultaniously
			over a 3 hours in 50 μl portions resulting suspension is
		·	further agitated for 10 hours to yild pale yellow colored
	•		solid powder
8	HRhCO (TPPTS)3, 50 mg,	Strontium chloride	Solution A and solution B were added to a suspension
	TPPTS 200 mg.	saturated solution in	of 2 gm bentonite in 10 ml water and resulting
	_		suspension is vigorously agitated to this suspension
	Dissolved in water 2 ml	water	solution A and solution B were added simultaniously
·			over a 3 hours in 50 µl portions resulting suspension is
			further agitated for 10 hours to yild pale yellow color
			solid powder
9	HRhCO (TPPTS)3, 50 mg,	Calcium chloride 500	Solution A and solution B were added to a suspension
	TPPTS 200 mg.	mg solution in 2 ml	of 2 gm bentonite in 10 ml water and resulting
	Dissolved in water 2 ml	water	suspension is vigorously agitated to this suspension
	,		solution A and solution B were added simultaniously
	•		over a 3 hours in 50 µl portions resulting suspension is
i l			further agitated for 10 hours to yild pale yellow color
	TINI CO CENTRALIA	D	solid powder
10	HRhCO (TPPTS)3, 50 mg,	Barium nitrate	Solution A and solution B were added to a suspension
	TPPTS 200 mg.	saturated solution in	of 2 gm charcoal in 10 ml water and resulting
	Dissolved in water 2 ml	water	suspension is vigorously agitated to this suspension
			solution A and solution B were added simultaneously
			over a 3 hours in 50 µl portions resulting suspension is
			further agitated for 2 hours to yield black colored solid
			powder

. 11	HRhCO (TPPTS)3, 50 mg, TPPTS 200 mg. Dissolved in water 2 ml	Strontium chloride saturated solution in water	Solution A and solution B were added to a suspension of 2 gm charcoal in 10 ml water and resulting suspension is vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 2 hours to yield black colored solid
			powder.
12	HRhCO (TPPTS)3, 50 mg,	Calcium chloride 500	Solution A and solution B were added to a suspension
	TPPTS 200 mg.	mg solution in 2 ml	of 2 gm charcoal in 10 ml water and resulting
ì	Dissolved in 2 ml water	water	suspension is vigorously agitated to this suspension
			solution A and solution B were added simultaneously
			over a 3 hours in 50 µl portions resulting suspension is
	,		further agitated for 2 hours to yield black colored solid
		·	powder
13	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	Solution A and solution B were added to a suspension
•	TPPTS 200 mg	saturated solution in	of 2 gm Davisil in 10 ml water and resulting suspension
	Dissolved in 2 ml water	2 ml water	is vigorously agitated to this suspension solution A and
			solution B were added simultaniously over a 3 hours in
			50 µl portions resulting suspension is further agitated
			for 10 hours to yild light brown colored solid powder.
14	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Strontium chloride	Solution A and solution B were added to a suspension
	TPPTS 200 mg	saturated solution in	of 2 gm Davisil in 10 ml water and resulting
·	Dissolved in 2 ml water.	2 ml water	suspension is vigorously agitated to this suspension
			solution A and solution B were added simultaniously
			over a 3 hours in 50 µl portions resulting suspension is
	·		further agitated for 10 hours to yield light brown
	D. GD (OI) GTDDTG), 50	7	colored solid powder. Solution A and solution B were added to a suspension
. 15	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate saturated solution in	
	TPPTS 200 mg Dissolved in 2 ml water	2 ml water	of 2 gm γ-alumina in 10 ml water and resulting suspension is vigorously agitated to this suspension
ļ	2 iii watei	2 im water	solution A and solution B were added simultaneously
	•		over a 3 hours in 50 µl portions resulting suspension is
l			further agitated for 10 hours to yield light brown
ĺ			colored solid powder.
16	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Strontium chloride	Solution A and solution B were added to a suspension
	TPPTS 200 mg Dissolved in	saturated solution in	of 2 gm y-alumina in 10 ml water and resulting
	2 ml water	2 ml water	suspension is vigorously agitated to this suspension
			solution A and solution B were added simultaneously
			over a 3 hours in 50 µl portions resulting suspension is
			further agitated for 10 hours to yield light brown
			colored solid powder.
17	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Strontium chloride	Solution A and solution B were added to a suspension
	TPPTS 200 mg	saturated solution in	of 2 gm \u03c4-alumina in 10 ml water and resulting
	Sodium polyvinylsulfonate	2 ml water	suspension is vigorously agitated to this suspension
	500 mg Dissolved in 2 ml		solution A and solution B were added simultaneously
	water		over a 3 hours in 50 µl portions resulting suspension is
			further agitated for 10 hours to yield light brown
			colored solid powder.
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18	Ru (H)(Cl)(TPPTS) 3 50 mg TPPTS 200 mg Sodium polyvinylsulfonate 500 mg Dissolved in 2 ml water	Barium nitrate saturated solution in 2 ml water	Solution A and solution B were added to a suspension of 2 gm γ -alumina in 10 ml water and resulting suspension is vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 μ l portions resulting suspension is further agitated for 10 hours to yield light brown colored solid powder.
19	Ru (H)(Cl)(TPPTS) 3 50 mg TPPTS 200 mg Sodium polyvinylsulfonate 500 mg Dissolved in 2 ml water	Barium nitrate saturated solution in 2 ml water	Solution A and solution B were added to a suspension of 2 gm titania in 10 ml water and resulting suspension is vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield light brown colored solid powder.
20	Ru (H)(Cl)(TPPTS) 3 50 mg TPPTS 200 mg Sodium polyvinylsulfonate 500 mg Dissolved in 2 ml water	Barium nitrate saturated solution in 2 ml water	Solution A and solution B were added to a suspension of 2 gm zirconia in 10 ml water and resulting suspension is vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield light brown colored solid powder.
21	Ru (H)(Cl)(TPPTS) 3 50 mg TPPTS 200 mg Sodium polyvinylsulfonate 500 mg Dissolved in 2 ml water	Barium nitrate saturated solution in 2 ml water	Solution A and solution B were added to a suspension of 2 gm activated charcoal in 10 ml water and resulting suspension is vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield black colored solid powder.
22	PdCl ₂ (TPPTS) ₂ 10 mg TPPTS 100 mg Poly acrylic acid sodium salt in 5 ml	Barium nitrate saturated solution 5 ml	Solution A and solution B were added to a suspension of 2 gm shreded asbestos roap in 20 ml water and resulting suspension is vigorously agitated to this suspension solution A and solution B was added s over a period of 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield yellow gray colored solid powder.
23	PdCl ₂ (TPPTS) ₂ 10 mg TPPTS 100 mg Poly acrylic acid sodium salt in 5 ml	Strontium chloride saturated solution 5 ml	Solution A and solution B were added to a suspension of 2 gm shreded asbestos roap in 20 ml water and resulting suspension is vigorously agitated to this suspension solution A and solution B was added over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield yellow gray colored solid powder.

24	PdCl ₂ (TPPTS) ₂ 10 mg	500 mg calcium	Solution A and solution B were added to a suspension
	TPPTS 100 mg	chloride in 5 ml	of 2 gm shreded asbestos roap in 20 ml water and
	Poly acrylic acid sodium salt	water.	resulting suspension is vigorously agitated to this
	in 5 ml		suspension solution A and solution B was added over a
		·	3 hours in 50 µl portions resulting suspension is further
			agitated for 10 hours to yield yellow gray colored solid
			powder.
25	PdAc ₂ BYPYDS 25 mg	Barium nitrate	Solution A and solution B were added to a suspension
	BYPYDS 100 mg	saturated solution	of 2 gm davisil in 20 ml water and resulting suspension
	Dissolved in 2 ml water	5ml	is vigorously agitated to this suspension solution A and
			solution B was added over a 3 hours in 50 µl portions
			resulting suspension is further agitated for 10 hours to
			yield pale orange colored solid powder.
26	PdAc2BYPYDS 25 mg	Strontium chloride	Solution A and solution B were added to a suspension
	BYPYDS 100 mg	saturated solution	of 2 gm davisil in 20 ml water and resulting suspension
	Dissolved in 2 ml water	5ml	is vigorously agitated to this suspension solution A and
	•		solution B was added over a 3 hours in 50 µl portions
			resulting suspension is further agitated for 10 hours to
			yield pale orange colored colored solid powder.
27	PdAc ₂ BYPYDS 25 mg	500 mg calcium	Solution A and solution B were added to a suspension
	BYPYDS 100 mg	chloride in 5 ml	of 2 gm davisil in 20 ml water and resulting suspension
	Dissolved in 2 ml water	water	is vigorously agitated to this suspension solution A and
			solution B was added over a 3 hours in 50 µl portions
	٠	•	resulting suspension is further agitated for 10 hours to
·			yield pale orange colored solid powder.
28	PdAc ₂ BYPYDS 25 mg	Barium nitrate	Solution A and solution B were added to a suspension
	BYPYDS 100 mg	saturated solution	of 2 gm bentonite in 20 ml water and resulting
	Dissolved in 2 ml water	5ml	suspension is vigorously agitated to this suspension
		· .	solution A and solution B was added over a 3 hours in
			50 µl portions resulting suspension is further agitated
	Dala and Calland S. S. S.	n-t	for 10 hours to yield light orange colored solid powder.
29	PdAc ₂ tri (o) tolyl phosphine	Barium nitrate	Solution A and solution B were added to a suspension
	trisulfonated 25 mg	saturated solution	of 2 gm bentonite in 20 ml water and resulting
	Tri (o) tolyl phosphine trisulfonated 100 mg	5ml	suspension is vigorously agitated to this suspension
·	Dissolved in 2 ml water		solution A and solution B was added over a 3 hours in 50 µl portions resulting suspension is further agitated
	Dissured in 2 iii walci		for 10 hours to yield pale yellow other colored solid
			powder.
30	PdAc ₂ tri (o) tolyl phosphine	Strontium chloride	Solution A and solution B were added to a suspension
30	trisulfonated 25 mg	saturated solution	of 2 gm bentonite in 20 ml water and resulting
	Tri (o)tolyl phosphine	5ml	suspension is vigorously agitated to this suspension
•	trisulfonated 100 mg	- 1111	solution A and solution B was added over a 3 hours in
·	Dissolved in 2 ml water		50 µl portions resulting suspension is further agitated
	water		for 10 hours to yield pale yellow other colored solid
			powder.

31	PdAc ₂ trio tolyl phosphine trisulfonated 25 mg Tri ortho tolyl phosphine trisulfonated 100 mg Dissolved in 2 ml water	Barium nitrate saturated solution 5ml	Solution A and solution B were added to a suspension of 2 gm alumina in 20 ml water and resulting suspension is vigorously agitated to this suspension solution A and solution B was added over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield pale yellow ocher colored solid powder.
32	PdAc ₂ tri ortho tolyl phosphine trisulfonated 25 mg Tri ortho tolyl phosphine trisulfonated 100 mg Dissolved in 2 ml water	Barium nitrate saturated solution 5ml	Solution A and solution B were added to a suspension of 2 gm charcoal in 20 ml water and resulting suspension is vigorously agitated to this suspension solution A and solution B was added over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield black colored solid powder.
33	NiCl ₂ .(TPPTS) ₂ 25 mg TPPTS 100 mg Sodium carboxy methyl cellulose 100 mg Dissolved in 2 ml	Saturated barium nitrate in 2 ml water	A suspension of 1 gm davisil in 5 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield solid powder almost white with blue tinge.
34	NiCl ₂ .(TPPTS) ₂ 25 mg TPPTS 100 mg Sodium carboxy methyl cellulose 100 mg Dissolved in 2 ml	Saturated barium nitrate in 2 ml water	A suspension of 1 gm alumina in 5 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield solid powder almost white with blue tinge.
35	NiCl ₂ .(TPPTS) ₂ 25 mg TPPTS 100 mg Sodium carboxy methyl cellulose 100 mg Dissolved in 2 ml	Saturated barium nitrate in 2 ml water	A suspension of 1 gm zirconia in 5 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield solid powder almost white with blue tinge.
36	NiCl ₂ .(TPPTS) ₂ 25 mg TPPTS 100 mg Sodium carboxy methyl cellulose 100 mg Dissolved in 2 ml	Saturated strontium chloride in 2 ml water	A suspension of 1 gm zirconia in 5 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield solid powder almost white with blue tinge.

37	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated strontium	A suspension of 1 gm titania in 5 ml water was formed
	TPPTS 100 mg	chloride in 2 ml	and resulting suspension was vigorously agitated to this
	Sodium carboxy methyl	water	suspension solution A and solution B were added
	cellulose 100 mg		simultaneously over a 3 hours in 50 µl portions
	Dissolved in 2 ml		resulting suspension is further agitated for 10 hours to
			yield solid powder almost white with blue tinge.
38	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated strontium	A suspension of 1 gm asbestos in 5 ml water was
	TPPTS 100 mg	chloride in 2 ml	formed and resulting suspension was vigorously
	Sodium carboxy methyl	water	agitated to this suspension solution A and solution B
	cellulose 100 mg		were added simultaneously over a 3 hours in 50 µl
	Dissolved in 2 ml		portions resulting suspension is further agitated for 10
			hours to yield gray colored solid powder.
39 .	(IrClCOD) 5 mg exchanged	Saturated strontium	A suspension of 1 gm davisil in 5 ml water was formed
	with TPPTS 100 mg.	chloride in 2 ml	and resulting suspension was vigorously agitated to this
	Poly acrylic acid sodium salt	water	suspension solution A and solution B were added
	100 mg		simultaneously over a 3 hours in 50 µl portions
·	In 2 ml water		resulting suspension is further agitated for 10 hours to
.			yield pale yellow colored solid powder.
40	(IrClCOD) 5 mg exchanged	Saturated strontium	A suspension of 1 gm keisulghur in 5 ml water was
	with TPPTS 100 mg.	chloride in 2 ml	formed and resulting suspension was vigorously
	Poly acrylic acid sodium salt	water	agitated to this suspension solution A and solution B
	100 mg		were added simultaneously over a 3 hours in 50 µl
	In 2 ml water	,	portions resulting suspension is further agitated for 10
			hours to yield pale yellow colored solid powder.
41	(IrClCOD) 5 mg exchanged	Saturated strontium	A suspension of 1 gm bentonite in 5 ml water was
	with TPPTS 100 mg.	chloride in 2 ml	formed and resulting suspension was vigorously
	Poly acrylic acid sodium salt	water	agitated to this suspension solution A and solution B
	100 mg		were added simultaneously over a 3 hours in 50 µl
	In 2 ml water		portions resulting suspension is further agitated for 10
			hours to yield pale yellow colored solid powder.
42	(RuCl ₂ COD) 5 mg exchanged	Saturated strontium	A suspension of 1 gm davisil in 5 ml water was formed
42	with diphenyl phosphino	chloride in 2 ml	and resulting suspension was vigorously agitated to this
	ethane tetrasulfonate 100 mg.	water	suspension solution A and solution B were added
	Poly acrylic acid sodium salt	Walci	simultaneously over a 3 hours in 50 µl portions
	-		resulting suspension is further agitated for 10 hours to
	100 mg		
4.5	In 2 ml water	Canada at a state of	yield pale yellow colored solid powder.
43	(RuCl ₂ COD) 5 mg exchanged	Saturated strontium	A suspension of 1 gm davisil in 5 ml water was formed
	with diphenyl phosphino	chloride in 2 ml	and resulting suspension was vigorously agitated to this
	ethane tetrasulfonate 100 mg.	water	suspension solution A and solution B were added
	Poly acrylic acid sodium salt		simultaneously over a 3 hours in 50 µl portions
	100 mg	·	resulting suspension is further agitated for 10 hours to
·	In 2 ml water		yield pale yellow colored solid powder.
44	(RuCl₂COD) 5 mg exchanged	500 mg calcium	A suspension of 1 gm davisil in 5 ml water was formed
	with diphenyl phosphino	chloride in 2 ml	and resulting suspension was vigorously agitated to this
	ethane tetrasulfonate 100 mg.	water	suspension solution A and solution B were added
1	Poly acrylic acid sodium salt		simultaneously over a 3 hours in 50 µl portions
J	100 mg		resulting suspension is further agitated for 10 hours to
. 1	In 2 ml water		yield pale yellow colored solid powder.

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45	Rh(COD)PF ₆ / S,S chiraphos tetrasulfonate 25 mg S,S chiraphos tetrasulfonate 25 mg Sodium alginate 100 mg dissolved in 2 ml water	Saturated strontium chloride solution 2 ml	A suspension of 1 gm davisil in 5 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield pale yellow colored solid powder.
46	Rh(COD)PF ₆ / S,S chiraphos tetrasulfonate 25 mg S,S chiraphos tetrasulfonate 25 mg Sodium alginate 100 mg dissolved in 2 ml water	Saturated barium nitrate solution 2 ml	A suspension of 1 gm davisil in 5 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield pale yellow colored solid powder.
47	Rh (COD) PF6/ S,S chiraphos tetrasulfonate 25 mg S,S chiraphos tetrasulfonate 25 mg Sodium alginate 100 mg dissolved in 2 ml water	Saturated barium nitrate solution 2 ml	A suspension of 1 gm alumina in 5 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield pale yellow colored solid powder.
48	Rh(COD)PF ₆ / S,S chiraphos tetrasulfonate 25 mg S,S chiraphos tetrasulfonate 25 mg Sodium alginate 100 mg dissolved in 2 ml water	Saturated barium nitrate solution 2 ml	A suspension of 1 gm titania in 5 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield pale yellow colored solid powder.
49	HRhCO (TPATS) 3 10 mg 100 mg TPATS Carboxy methyl cellulose sodium 100 mg in 1 ml water	500 mg Calcium chloride solution in water 5 ml	A suspension of 1 gm titania in 5 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield pale yellow colored solid powder.
50	HRhCO (TPATS); 10 mg 100 mg TPATS carboxy methyl cellulose sodium 100 mg in 1 ml water	Strontium chloride saturated solution in water 5 ml	A suspension of 1 gm alumina in 5 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield pale yellow colored solid powder.
51	HRhCO (TPATS); 10 mg 100 mg TPATS carboxy methyl cellulose sodium 100 mg in 1 ml water	Barium nitrate saturated solution in water 5 ml	A suspension of 1 gm bentonite in 5 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield pale yellow colored solid powder.

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52	HRhCO (TPATS), 10 mg 100 mg TPATS Carboxy methyl cellulose sodium 100 mg in 1 ml water	Strontium chloride saturated solution in water 5 ml	A suspension of 1 gm titania in 5 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield pale yellow colored solid powder.
53	HRhCO (TPATS) ₃ 10 mg 100 mg TPATS Carboxy methyl cellulose sodium 100 mg in 1 ml water	Strontium chloride saturated solution in water 5 ml	A suspension of 1 gm Davisil in 5 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield pale yellow colored solid powder.
54	HRhCO (BISBIS) 50 mg BISBIS 200 mg 200 mg sodium sulfate Dissolved in 2 ml water	Saturated barium nitrate solution is 5 ml water	A suspension of 2 gm Davisil in 5 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield pale yellow colored solid powder.
55	HRhCO (BISBIS) 50 mg BISBIS 200 mg 200 mg polyvinyl sulfonic acid dissolved in 2 ml water	1 g calcium chloride solution In 5 ml water	A suspension of 2 gm Davisil in 10 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield pale yellow colored solid powder.
56	HRhCO(BISBIS) 50 mg BISBIS 200 mg 200 mg polyacrylic acid sodium salt dissolved in 2 ml water	Saturated barium nitrate solution is 5 ml water	A suspension of 2 gm titania in 10 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield pale yellow colored solid powder.7
57	HRhCO(BISBIS) 50 mg BISBIS 200 mg 200 mg polyvinyl sulfonic acid dissolved in 2 ml water	Saturated strontium chloride solution is 5 ml water	A suspension of 2 gm alumina in 10 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield pale yellow colored solid powder.
58	HRhCO(BISBIS) 50 mg BISBIS 200 mg 200 mg polyvinyl sulfonic acid dissolved in 2 ml water	Saturated barium nitrate solution is 5 ml water	A suspension of 2 gm bentonite in 10 ml water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added simultaneously over a 3 hours in 50 µl portions resulting suspension is further agitated for 10 hours to yield pale yellow colored solid powder.

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59	9.	HRhCO(BISBIS) 50 mg	Saturated barium	A suspension of 2 gm Davisil in 10 ml water was
		BISBIS 200 mg	nitrate solution is 5	formed and resulting suspension was vigorously
		200 mg polyvinyl sulfonic	ml water	agitated to this suspension solution A and solution B
		acid dissolved in 2 ml water		were added simultaneously over a 3 hours in 50 μ l
	•			portions resulting suspension is further agitated for 10
			·	hours to yield pale yellow colored solid powder.
60	0	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	A suspension of 2 gm of davisil in 10 ml butane diol
		TPPTS 100 mg	barium nitrate 5 ml	was formed and resulting suspension was vigorously
		100 mg sodium alginate	:	agitated to this suspension solution A was added over a
•		dissolved in 2 ml water		period of 2 hours and further agitated for 5 hours
				solution B was then added in portions of 50µl over a
Į.				period of 3 hours resulting suspension is further
1				agitated for 24 hours to yield gray colored solid
		•		powder.
<u></u>	-	Deal (mppmas) 50	Commend colusion of	
61	•	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	A suspension of 2 gm of y-alumina in 10 ml butane
		TPPTS 100 mg	barium nitrate 5 ml	diol was formed and resulting suspension was
	.	100 mg oxalic acid sodium		vigorously agitated to this suspension solution Awas
	. 1	salt.		added over aperiod of 2 hours and further agitated for 5
l		Dissolved in 2 ml water		hours solution B was then added in portions of 50µl
				over a period of 3 hours resulting suspension is further
				agitated for 24 hours to yield pale yellow colored solid
,				powder.
62	2	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	A suspension of 2 gm of davisil in 10 ml ethylene
· .		TPPTS 100 mg	strontium chloride 5	glycol was formed and resulting suspension was
		100 mg citric acid	ml	vigorously agitated to this suspension solution A was
1		Dissolved in 2 ml water		added over aperiod of 2 hours and further agitated for 5
1				hours solution B was then added in portions of 50µl
				over a period of 3 hours resulting suspension is further
				agitated for 24 hours to yield pale yellow colored solid
	·		·	powder.
63	3	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	A suspension of 2 gm of davisil in 10 ml butane diol
1		TPPTS 100 mg	barium nitrate 5 ml	was formed and resulting suspension was vigorously
		100 mg polyacrylic acid	•	agitated to this suspension solution A was added over
		sodium salt.		aperiod of 2 hours and further agitated for 5 hours
		Dissolved in 2 ml water		solution B was then added in portions of 50µl over a
				period of 3 hours resulting suspension is further
		•		agitated for 24 hours to yield pale yellow colored solid
				powder.
64	4	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	A suspension of 2 gm of shreded asbestos roap in 10
`		TPPTS 300 mg	barium nitrate 5 ml	ml butane diol was formed and resulting suspension
		Dissolved in 2 ml water		was vigorously agitated to this suspension solution
				Awas added over aperiod of 2 hours and further
	ļ	,		agitated for 5 hours solution B was then added in
				portions of 50µl over a period of 3 hours resulting
1				• • • • • • • • • • • • • • • • • • •
I			l	suspension is further agitated for 24 hours to yield gray
			ľ	colored solid powder.

65	Cobalt N, N'ethylene bis (salicyldiamine) 5-sulfonato sodium 100 mg. Sodium phosphate. 500 mg. In 5 ml water	Saturated barium nitrate solution in water 5ml	A suspension of 2 gm davisil in 10 ml tetrahydrofuran 50 % in water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added at simultaniously in portions of 50µl over a period of 3 hours resulting suspension is further agitated for 24 hours to yield pale brown colored solid powder.
66	Cobalt N, N'ethylene bis (salicyldiamine) 5-sulfonato sodium 100 mg. Sodium silicate 500 mg. In 5 ml water	Saturated barium nitrate solution in water 5ml	A suspension of 2 gm alumina in 10 ml tetrahydrofuran 50 % in water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added at simultaniously in portions of 50µl over a period of 3 hours resulting suspension is further agitated for 24 hours to yield pale brown colored solid powder.
67	Cobalt N, N'ethylene bis (salicyldiamine) 5-sulfonato sodium 100 mg. Polyvinyl sulfonate sodium. 500 mg. In 5 ml water	Saturated barium nitrate solution in water 5ml	A suspension of 2 gm titania in 10 ml tetrahydrofuran 50 % in water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added at simultaniously in portions of 50µl over a period of 3 hours resulting suspension is further agitated for 24 hours to yield pale brown colored solid powder.
68	Cobalt N, N'ethylene bis (salicyldiamine) 5-sulfonato sodium 100 mg. Polyvinyl sulfonate sodium. 500 mg. In 5 ml water	Saturated barium nitrate solution in water 5ml	A suspension of 2 gm zirconia asbesto rope in 10 ml tetrahydrofuran 50 % in water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added at simultaniously in portions of 50µl over a period of 3 hours resulting suspension is further agitated for 24 hours to yield pale brown colored solid powder.
69	Cobalt N, N'ethylene bis (salicyldiamine) 5-sulfonato sodium 100 mg. Polyvinyl sulfonate sodium. 500 mg. In 5 ml water	2g calcium chloride solution in water 5ml	A suspension of 2 gm shreded asbesto rope in 10 ml tetrahydrofuran 50 % in water was formed and resulting suspension was vigorously agitated to this suspension solution A and solution B were added at simultaniously in portions of 50µl over a period of 3 hours resulting suspension is further agitated for 24 hours to yield pale brown colored solid powder.
70	Cobalt (II), 4, 4', 4",4"'- tetrasulfopthalocynine oxygen adduct. 500 mg And 500 mg sodium sodium poly vinyl sulfonate in 5 ml water	Saturated strontium chloride in 5 ml water	A suspension of 2 gm shreded asbesto rope in 10 ml methanol was formed and resulting suspension was vigorously agitated to this suspension solution A was added and stirred for 15 min and solution B was added at once resulting suspension is further agitated for 3 hours to yield steel gray colored solid powder.

71	Cobalt (II), 4, 4', 4",4"'-	Saturated barium	A suspension of 2 gm keisulghur in 10 ml methanol
	tetrasulfopthalocynine. 500	nitrate in 5 ml water	was formed and resulting suspension was vigorously
	mg		agitated to this suspension solution A was added and
	And 500 mg sodium		stirred for 15 min and solution B was added at once
	phosphate in 5 ml water		resulting suspension is further agitated for 3 hours to
	1.		yield light blue colored solid powder.
72	Cobalt (II), 4, 4', 4",4"'-	Saturated strontium	A suspension of 2 gm keisulghur in 10 ml methanol
'2	tetrasulfopthalocynine . 500	chloride in 5 ml	was formed and resulting suspension was vigorously
	· · ·	water	agitated to this suspension solution A was added and
	mg And 500 mg sodium	water .	stirred for 15 min and solution B was added at once
	1		resulting suspension is further agitated for 3 hours to
	phosphate in 5 ml water		
		·	yield light blue colored solid powder.
73	Cobalt (II), 4, 4', 4",4"'-	500mg. CaCl ₂ in 5 ml	A suspension of 2 gm keisulghur in 10 ml methanol
	tetrasulfopthalocynine. 500	water	was formed and resulting suspension was vigorously
	mg	٠.	agitated to this suspension solution A was added and
	And 500 mg sodium		stirred for 15 min and solution B was added at once
	phosphate in 5 ml water		resulting suspension is further agitated for 3 hours to
•		٠	yield light blue colored solid powder.
74	Copper (II), 4, 4', 4",4"'-	500mg. CaCl ₂ in 5 ml	A suspension of 2 gm keisulghur in 10 ml methanol
	tetrasulfopthalocynine. 500	water	was formed and resulting suspension was vigorously
	mg	,	agitated to this suspension solution A was added and
<u> </u>	And 500 mg sodium sulfate in		stirred for 15 min and solution B was added at once
	5 ml water		resulting suspension is further agitated for 3 hours to
	V	•	yield light blue colored solid powder.
75	Copper (II), 4, 4', 4'',4'''-	Saturated strontium	A suspension of 2 gm keisulghur in 10 ml methanol
/3	tetrasulfopthalocynine. 500	chloride in 5 ml	was formed and resulting suspension was vigorously
		water	agitated to this suspension solution A was added and
	mg	water	stirred for 15 min and solution B was added at once
l	And 500 mg sodium silicate		resulting suspension is further agitated for 3 hours to
	in 5 ml water		
·			yield light blue colored solid powder.
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76	Copper (II), 4, 4', 4'',4'''-	Saturated barium	A suspension of 2 gm keisulghur in 10 ml methanol
	tetrasulfopthalocynine. 500	nitrate in 5 ml water	was formed and resulting suspension was vigorously
	mg		agitated to this suspension solution A was added and
	And 500 mg sodium silicate		stirred for 15 min and solution B was added at once
	in 5 ml water		resulting suspension is further agitated for 3 hours to
		,	yield light blue colored solid powder.
			·
			
77	Copper (II), 4, 4', 4'',4'''-	Saturated barium	A suspension of 2 gm bentonite in 10 ml methanol was
- •	tetrasulfopthalocynine . 500	nitrate in 5 ml water	formed and resulting suspension was vigorously
, ,	mg		agitated to this suspension solution A was added and
l l			afrance to mis suspension sommon is was added and
	•		stirred for 15 min and colution R una added at anna
	And 500 mg sodium silicate		stirred for 15 min and solution B was added at once
	•		resulting suspension is further agitated for 3 hours to
·	And 500 mg sodium silicate	· · · · · ·	

78	Copper (II), 4, 4', 4'',4'''-	Saturated strontiun	A suspension of 2 gm bentonite in 10 ml methanol was
٠.	tetrasulfopthalocynine. 500	chloride in 5 ml	formed and resulting suspension was vigorously
	mg .	water	agitated to this suspension solution A was added and
•	And 500 mg sodium silicate	,	stirred for 15 min and solution B was added at once
	in 5 ml water		resulting suspension is further agitated for 3 hours to
			yield light blue colored solid powder.
			•
79	Manganese(II), 4, 4', 4'',4'''-	Saturated strontiun	A suspension of 2 gm Davisil in 10 ml methanol was
	tetrasulfopthalocynine. 500	chloride in 5 ml	formed and resulting suspension was vigorously
	mg	water	agitated to this suspension solution A was added and
	And 500 mg sodium silicate		stirred for 15 min and solution B was added at once
	in 5 ml water		resulting suspension is further agitated for 3 hours to
			yield light blue colored solid powder.
	•		
80	Manganese(II), 4, 4', 4'',4'''-	Saturated barium	A suspension of 2 gm Davisil in 10 ml methanol was
	tetrasulfopthalocynine. 500	nitrate in 5 ml water	formed and resulting suspension was vigorously
	ing		agitated to this suspension solution A was added and
	And 500 mg sodium silicate		stirred for 15 min and solution B was added at once
	in 5 ml water		resulting suspension is further agitated for 3 hours to
	III J III Water		yield light blue colored solid powder.
			yield light blue colored solid powder.
	Manganese(II), 4, 4', 4'',4'''-	Saturated barium	A suspension of 2 gm γ-alumina in 10 ml methanol
81	· ·		•
•		nitrate in 5 ml water	was formed and resulting suspension was vigorously
·	mg		agitated to this suspension solution A was added and
·	And 500 mg sodium silicate		stirred for 15 min and solution B was added at once
	in 5 ml water		resulting suspension is further agitated for 3 hours to
·			yield light blue colored solid powder.
82	Manganese(II), 4, 4', 4'',4'''-	Saturated barium nitrate in 5 ml water	A suspension of 2 gm γ-alumina in 10 ml methanol
	tetrasulfopthalocynine 500	nitrate in 5 mi water	was formed and resulting suspension was vigorously
·	mg		agitated to this suspension solution A was added and
	And 500 mg sodium	,	stirred for 15 min and solution B was added at once
	polyvinyl sulfonate in 5 ml		resulting suspension is further agitated for 3 hours to
	water		yield light blue colored solid powder.
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83	Iron (III), 4, 4', 4'',4'''-	Saturated strontium	A suspension of 2 gm Davisil in 10 ml methanol was
	tetrasulfopthalocynine oxygen	chloride in 5 ml	formed and resulting suspension was vigorously
	adduct. 500 mg	water	agitated to this suspension solution A was added and
. ,	And 500 mg sodium sulfate in		stirred for 15 min and solution B was added at once
	5 ml water		resulting suspension is further agitated for 3 hours to
		,	yield light blue colored solid powder.
84	Iron (III), 4, 4', 4'',4'''-	Saturated barium	A suspension of 2 gm Davisil in 10 ml methanol was
	tetrasulfopthalocynine oxygen	nitrate in water 5 ml	formed and resulting suspension was vigorously
	adduct. 500 mg		agitated to this suspension solution A was added and
	And 500 mg sodium sulfate in		stirred for 15 min and solution B was added at once
· .	5 ml water		resulting suspension is further agitated for 3 hours to
			yield light blue colored solid powder.
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Examples 85 to 168

Preparation of catalytic formulation by deposition precipitation

The following examples illustrate one of the procedures for the preparation of the catalytic formulation of the invention in accordance with the method of formulation known as co precipitation near the surface of the solid support.

The general procedure for the preparation of heterogeneous catalytic formulation is described herein as making of a solution of anionically charged catalytic entity, catalytically inert anionic additive (termed as solution A) and solution of group II A metal ions (termed as solution B). The specified amount of support pretreated as described in earlier is impregnated with solution A by wetting solids with solution followed by evaporation to obtain dry solid support bearing anionic components of solution A. this solid powder is gradually added to the solution B over a specified period of time. Resulting suspension is further agitated for specified time. Suspension was centrifuged and solids were repeatedly washed with water and dried in vacuum. Dry powder was stored under argon in gas tight vessel. These solid catalytic formulations can be used for appropriate reactions depending upon catalytically active entity incorporated in it.

Note1: solution A is prepared by dissolving anionic components including anionic complex and additives to make homogeneous solution in degassed solvents. And resulting solution is also degassed by purging argon.

Note 2: solution B is prepared by dissolving dissolving group IIA metal salts. Solution was degassed prior to use

Note 3: the impregnation of solution on solid support is carried out by wetting solids with solution A and evaporating in vacuum at 50 0 C unless stated

Note 4: addition of impregnated solids with components of A to solution B is carried out at ambient temperature unless stated.

Example	Solution A	Solution B	Procedure
85	HRhCO (TPPTS); 50 mg, TPPTS 200 mg. Dissolved in water 2 ml	Saturated barium nitrate in water 2 ml	2 gm Davisil was wetted with 100 μl portion of solution A and evaporated under vaccume 10 mm Hg with simultanious tumbling remining solution A was added in 100 μl fractions and solid was isolated (moisture content -20 %) this powder was added to solution B in equal fractions over a period of 2 hours and suspension was further agitated for 10 hours to yild pale yellow colored solid powder.

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86	HRhCO (TPPTS) 3 50 mg,	Saturated strontium	2 gm Davisil was wetted with 100 μl portion of
	TPPTS 200 mg.	chloride in water 2ml	solution A and evaporated under vaccume 10 mm Hg
	Dissolved in water 2 ml		with simultanious tumbling remining solution A was
			added in 100 µl fractions and solid was isolated
			(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours to yild
			pale yellow colored solid powder.
87	HRhCO (TPPTS)3, 50 mg,	500 mg of calcium	2 gm Davisil was wetted with 100 μl portion of
	TPPTS 200 mg.	chloride in 2 ml	solution A and evaporated under vaccume 10 mm Hg
	Dissolved in water 2 ml	water	with simultanious tumbling remining solution A was
			added in 100 µl fractions and solid was isolated
			(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
,			and suspension was further agitated for 10 hours to yild
		·	pale yellow colored solid powder.
88	HRhCO(TPPTS)3, 50 mg,	Barium nitrate	2 gm γ-alumina was wetted with 100 μl portion of
	TPPTS 200 mg.	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
· ·	Dissolved in water 2 ml	water	with simultanious tumbling remining solution A was
			added in 100 µl fractions and solid was isolated
		•	(moisture content -20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours to yild
			pale yellow colored solid powder.
89	HRhCO(TPPTS)3, 50 mg,	Strontium chloride	2 gm γ-alumina was wetted with 100 μl portion of
	TPPTS 200 mg.	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	Dissolved in water 2 ml	water 2ml	with simultanious tumbling remining solution A was
			added in 100 µl fractions and solid was isolated
• •			(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours to yild
		•	pale yellow colored solid powder.
90	HRhCO(TPPTS)3, 50 mg,	Calcium chloride 500	2 gm γ-alumina was wetted with 100 μl portion of
30	TPPTS 200 mg.	mg solution in 2 ml	
	Dissolved in water 2 ml	water	solution A and evaporated under vaccume 10 mm Hg
	Dissolved in water 2 iiii	watt.	with simultanious tumbling remining solution A was
			added in 100 µl fractions and solid was isolated
, ,			(moisture content ~20 %) this powder was added to
. 1			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours,
			filtered to yild pale yellow colored solid powder.
91	HRhCO(TPPTS)3, 50 mg,	Barium nitrate	2 gm bentonite was wetted with 100 μl portion of
	TPPTS 200 mg.	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
·	Dissolved in water 2 ml	water 2 ml	with simultanious tumbling remining solution A was
·			added in 100 µl fractions and solid was isolated
			(moisture content ~20 %)this powder was added to
			solution B in equal fractions over a period of 2 hours
	•		and suspension was further agitated for 10 hours to yild
			pale yellow colored solid powder.

92	HRhCO(TPPTS)3, 50 mg,	Strontium chloride	2 gm bentonite was wetted with 100 μl portion of
	TPPTS 200 mg.	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	Dissolved in water 2 ml	water	with simultanious tumbling remining solution A was
			added in 100 µl fractions and solid was isolated
			(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours to yild
٠			pale yellow colored solid powder.
93	HRhCO(TPPTS)3, 50 mg,	Calcium chloride 500	2 gm bentonite was wetted with 100 µl portion of
	TPPTS 200 mg.	mg solution in 2 ml	solution A and evaporated under vaccume 10 mm Hg
	Dissolved in water 2 ml	water	with simultanious tumbling remining solution A was
		•	added in 100 µl fractions and solid was isolated
		·	(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours to yild
		. :	pale yellow colored solid powder.
94	HRhCO(TPPTS)3, 50 mg,	Barium nitrate	2 gm charcoal was wetted with 100 µl portion of
	TPPTS 200 mg.	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	Dissolved in water 2 ml	water	with simultanious tumbling remining solution A was
	Dissolved in water 2 in	*********	added in 100 µl fractions and solid was isolated
			(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
		,	and suspension was further agitated for 10 hours to yild
			black colored solid powder.
	TYPI-COCTEDERS 50	Strontium chloride	2 gm charcoal was wetted with 100 µl portion of
95	HRhCO(TPPTS)3, 50 mg,	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
İ	TPPTS 200 mg. Dissolved in water 2 ml	water	with simultanious tumbling remining solution A was
	Dissolved in water 2 im	Water	added in 100 µl fractions and solid was isolated
	·		(moisture content ~20 %) this powder was added to
			and suspension was further agitated for 10 hours to yild
			ł ·
			black colored solid powder.
96	HRhCO(TPPTS)3, 50 mg,	Calcium chloride 500	2 gm charcoal was wetted with 100 µl portion of
	TPPTS 200 mg.	mg solution in 2 ml	solution A and evaporated under vaccume 10 mm Hg
	Dissolved in 2 ml water	water	with simultanious tumbling remining solution A was
			added in 100 µl fractions and solid was isolated
	·		(moisture content ~20 %)this powder was added to
			solution B in equal fractions over a period of 2 hours
		·	and suspension was further agitated for 10 hours to yild
		·	black colored solid powder.
97	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	2 gm Davisil was wetted with 100 μl portion of
	TPPTS 200 mg	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	Dissolved in 2 ml water	2 ml water	with simultanious tumbling remining solution A was
		·	added in 100 µl fractions and solid was isolated
			(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
•			filtured to yield light brown colored solid powder
			<u> </u>

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98	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Strontium chloride	2 gm Davisil was wetted with 100 µl portion of
•	TPPTS 200 mg	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	Dissolved in 2 ml water.	2 ml water	with simultanious tumbling remining solution A was
			added in 100 µl fractions and solid was isolated
			(moisture content ~20 %) this powder was added to
,			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
	•		filtured to yield light brown colored solid powder.
•			·
99	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	2 gm γ-alumina was wetted with 100 μl portion of
	TPPTS 200 mg Dissolved in	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	2 ml water	2 ml water	with simultanious tumbling remining solution A was
			added in 100 µl fractions and solid was isolated
		,	(moisture content ~20 %) this powder was added to
	•		solution B in equal fractions over a period of 2 hours
٠			and suspension was further agitated for 10 hours
			filtered to yield light brown colored solid powder.
100	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Strontium chloride	2 gm γ-alumina was wetted with 100 μl portion of
	TPPTS 200 mg Dissolved in	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
•	2 ml water	2 ml water	with simultanious tumbling remining solution A was
,			added in 100 µl fractions and solid was isolated
	•		(moisture content ~20 %) this powder was added to
	• •	•	solution B in equal fractions over a period of 2 hours
		-	and suspension was further agitated for 10 hours
	·		filtered to yield light brown colored solid powder.
101	Ru(H)(Cl)(TPPTS) ₂ 50 mg	Strontium chloride	2 gm γ-alumina was wetted with 100 μl portion of
	TPPTS 200 mg	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	Sodium polyvinylsulfonate	2 ml water	with simultanious tumbling remining solution A was
	500 mg Dissolved in 2 ml		added in 100 µl fractions and solid was isolated
<i>*</i>	water		(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
	•		and suspension was further agitated for 10 hours
			filtered to yield light brown colored solid powder.
102	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	2 gm γ-alumina was wetted with 100 μl portion of
102	TPPTS 200 mg	saturated solution in	
	Sodium polyvinylsulfonate	2 ml water	solution A and evaporated under vaccume 10 mm Hg
,	500 mg Dissolved in 2 ml	~ III Water	with simultanious tumbling remining solution A was
	water		added in 100 µl fractions and solid was isolated
	mattel		(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
,		·	and suspension was further agitated for 10 hours
			filtered to yield light brown colored solid powder
103	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	2 gm titania was wetted with 100 μl portion of solution
	TPPTS 200 mg	saturated solution in	A and evaporated under vaccume 10 mm Hg with
	Sodium polyvinylsulfonate	2 ml water	simultanious tumbling remining solution A was added
	500 mg Dissolved in 2 ml		in 100 µl fractions and solid was isolated (moisture
·	water		content ~20 %) this powder was added to solution B in
			equal fractions over a period of 2 hours and suspension
			was further agitated for 10 hours filtered to yield light
		.	brown colored solid powder.
-			

104	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	2 gm zirconia was wetted with 100 μl portion of
	TPPTS 200 mg	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	Sodium polyvinylsulfonate	2 ml water	with simultanious tumbling remining solution A was
	500 mg Dissolved in 2 ml		added in 100 µl fractions and solid was isolated
	water	`.	(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
			filtered to yield light brown colored solid powder
105	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	2 gm activated charcoal was wetted with 100 µl portion
	TPPTS 200 mg	saturated solution in	of solution A and evaporated under vaccume 10 mm
•	Sodium polyvinylsulfonate	2 ml water	Hg with simultanious tumbling remining solution A
	500 mg Dissolved in 2 ml		was added in 100 µl fractions and solid was isolated
	water		(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
		*	filtered to yield black colored solid powder.
106	PdCl ₂ (TPPTS) ₂ 10 mg	Barium nitrate	2 gm shreaded asbestos roap was wetted with 100 μl
100	TPPTS 100 mg	saturated solution 5	portion of solution A and evaporated under vaccume 10
	Poly acrylic acid sodium salt	ml	mm Hg with simultanious tumbling remining solution
	in 5 ml		A was added in 100 µl fractions and solid was isolated
			(moisture content ~20 %) this powder was added to
	·	•	solution B in equal fractions over a period of 2 hours
	<u> </u>	·	and suspension was further agitated for 10 hours
			filtered to yield yellow gray colored solid powder.
	PdCl ₂ (TPPTS) ₂ 10 mg	Strontium chloride	2 gm shreaded asbestos roap was wetted with 100 μl
107	TPPTS 100 mg	saturated solution 5	portion of solution A and evaporated under vaccume 10
	Poly acrylic acid sodium salt	ml	mm Hg with simultanious tumbling remining solution
	in 5 ml		A was added in 100 µl fractions and solid was isolated
	111.5 till		(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
		·	filtered to yield yellow gray colored solid powder.
108	PdCl ₂ (TPPTS) ₂ 10 mg	500 mg calcium	2 gm shreaded asbestos roap was wetted with 100 μl
•	TPPTS 100 mg	chloride in 5 ml	portion of solution A and evaporated under vaccume 10
	Poly acrylic acid sodium salt	water.	mm Hg with simultanious tumbling remining solution
	in 5 ml		A was added in 100 µl fractions and solid was isolated
			(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
			filtered to yield yellow gray colored solid powder.
109	PdAc ₂ BYPYDS 25 mg	Barium nitrate	2 gm davisil was wetted with 100 μl portion of solution
	BYPYDS 100 mg	saturated solution	A and evaporated under vaccume 10 mm Hg with
	Dissolved in 2 ml water	5ml	simultanious tumbling remining solution A was added
			in 100 µl fractions and solid was isolated (moisture
			content ~20 %) this powder was added to solution B in
	•	ł	
			equal fractions over a period of 2 hours and suspension
			equal fractions over a period of 2 hours and suspension was further agitated for 10 hours filtered to yield loght

PdAc ₂ BYPYDS 25 mg Strontium chloride 2 gm davisil was wetted with 100 BYPYDS 100 mg saturated solution A and evaporated under vaccum	µl portion of solution
RYPYDS 100 mg I saturated solution A and evanorated under vaccing	-
Dissolved in 2 ml water 5 ml simultanious tumbling remining s	olution A was added
in 100 µl fractions and solid w	as isolated (moisture
content ~20 %) this powder was a	dded to solution B in
equal fractions over a period of 2	hours and suspension
was further agitated for 10 hours	filtered to yield loght
orange colored solid powder.	
111 PdAc ₂ BYPYDS 25 mg 500 mg calcium 2 gm davisil was wetted with 100	μl portion of solution
BYPYDS 100 mg chloride in 5 ml A and evaporated under vaccum	ne 10 mm Hg with
Dissolved in 2 ml water water simultanious tumbling remining s	olution A was added
in 100 µl fractions and solid w	
content ~20 %) this powder was a	•
equal fractions over a period of 2	•
was further agitated for 10 hours	
orange colored solid powder.	. , , , , , , , , , , , , , , , , , , ,
	h 100 µl nortion of
Dissolved in 2 ml water 5 ml with simultanious tumbling remi	
added in 100 µl fractions and	•
(moisture content ~20 %) this p	
solution B in equal fractions over	
and suspension was further ag	itated for 10 hours
filtered to yield loght orange color	ed solid powder.
113 PdAc ₂ trio tolyl phosphine Barium nitrate 2 gm bentonite was wetted wit	h 100 µl portion of
trisulfonated 25 mg saturated solution solution A and evaporated under	vaccume 10 mm Hg
trio tolyl phosphine 5ml with simultanious tumbling remi	ning solution A was
trisulfonated 100 mg added in 100 µl fractions and	solid was isolated
Dissolved in 2 ml water (moisture content ~20 %) this p	owder was added to
solution B in equal fractions over	a period of 2 hours
and suspension was further ag	itated for 10 hours
filtered to yield pale brown colored	d solid powder.
114 PdAc ₂ trio tolyl phosphine Strontium chloride 2 gm bentonite was wetted with	h 100 µl portion of
trisulfonated 25 mg saturated solution solution A and evaporated under	
trio tolyl phosphine 5ml with simultanious tumbling remi	
trisulfonated 100 mg added in 100 µl fractions and	solid was isolated
Dissolved in 2 ml water (moisture content ~20 %) this p	
solution B in equal fractions over	
and suspension was further ag	-
filtered to yield pale brown colored	
	
	•
trio tolyl phosphine 5ml with simultanious tumbling remin	_
trisulfonated 100 mg added in 100 µl fractions and	
Dissolved in 2 ml water (moisture content ~20 %) this p	
solution B in equal fractions over	-
and suspension was further ag	
, , , , , , , , , , , , , , , , , , ,	i solid powder.

			·
116	PdAc ₂ trio tolyl phosphine	Barium nitrate	2 gm charcoal was wetted with 100 μl portion of
	trisulfonated 25 mg	saturated solution	solution A and evaporated under vaccume 10 mm Hg
	trio tolyl phosphine	5ml	with simultanious tumbling remining solution A was
i.	trisulfonated 100 mg		added in 100 µl fractions and solid was isolated
	Dissolved in 2 ml water		(moisture content -20 %) this powder was added to
		•	solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 24 hours
			filtered to yield black colored solid powder.
117	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated barium	1 gm davisil was wetted with 100 µl portion of solution
117	TPPTS 100 mg	nitrate in 2 ml water	A and evaporated under vaccume 10 mm Hg with
	Sodium carboxy methyl	muace in 2 iii water	
			simultanious tumbling remining solution A was added
	cellulose 100 mg		in 100 µl fractions and solid was isolated (moisture
	Dissolved in 2 ml		content ~20 %) this powder was added to solution B in
			equal fractions over a period of 2 hours and suspension
		•	was further agitated for 10 hours filtered to yield pale
		· . ·	blue colored solid powder.
118	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated barium	1 gm alumina was wetted with 100 μl portion of
	TPPTS 100 mg	nitrate in 2 ml water	solution A and evaporated under vaccume 10 mm Hg
	Sodium carboxy methyl	,	with simultanious tumbling remining solution A was
	cellulose 100 mg		added in 100 µl fractions and solid was isolated
	Dissolved in 2 ml		(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
·			filtered to yield pale blue colored solid powder.
119	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated barium	1 gm zirconia was wetted with 100 μl portion of
•	TPPTS 100 mg	nitrate in 2 ml water	solution A and evaporated under vaccume 10 mm Hg
,	Sodium carboxy methyl	·	with simultanious tumbling remining solution A was
	cellulose 100 mg	•	added in 100 µl fractions and solid was isolated
	Dissolved in 2 ml		(moisture content ~20 %) this powder was added to
.*	,		solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
			filtered to yield pale blue colored solid powder.
120	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated strontium	1 gm zirconia was wetted with 100 µl portion of
120			
	TPPTS 100 mg		solution A and evaporated under vaccume 10 mm Hg
	Sodium carboxy methyl	water	with simultanious tumbling remining solution A was
	cellulose 100 mg		added in 100 µl fractions and solid was isolated
,	Dissolved in 2 ml		(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
•			and suspension was further agitated for 10 hours
			filtered to yield pale blue colored solid powder.
121	NiCl₂.(TPPTS)₂ 25 mg	Saturated strontium	1 gm titania was wetted with 100 μl portion of solution
	TPPTS 100 mg	chloride in 2 ml	A and evaporated under vaccume 10 mm Hg with
	Sodium carboxy methyl	water	simultanious tumbling remining solution A was added
	cellulose 100 mg		in 100 µl fractions and solid was isolated (moisture
	Dissolved in 2 ml		content ~20 %) this powder was added to solution B in
			equal fractions over a period of 2 hours and suspension
			was further agitated for 10 hours filtered to yield pale
,			blue colored solid powder.
			<u> </u>

122	NICL CEPPTS: 25 mg	Saturated strontium	1 gm asbestos was wetted with 100 μl portion of
122.	NiCl ₂ .(TPPTS) ₂ 25 mg	chloride in 2 ml	solution A and evaporated under vaccume 10 mm Hg
	TPPTS 100 mg		· ·
	Sodium carboxy methyl	water	with simultanious tumbling remining solution A was
	cellulose 100 mg	, ·	added in 100 µl fractions and solid was isolated
•	Dissolved in 2 ml		(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
		·	and suspension was further agitated for 10 hours
			filtered to yield pale blue colored solid powder.
123.	(IrClCOD) 5 mg exchanged	Saturated strontium	1 gm davisil was wetted with 100 μl portion of solution
	with TPPTS 100 mg.	chloride in 2 ml	A and evaporated under vaccume 10 mm Hg with
	Poly acrylic acid sodium salt	water	simultanious tumbling remining solution A was added
•	100 mg	·	in 100 µl fractions and solid was isolated (moisture
	In 2 ml water	<u> </u>	content ~20 %) this powder was added to solution B in
			equal fractions over a period of 2 hours and suspension
			was further agitated for 10 hours filtered to yield pale
		2	yellow colored solid powder.
124.	(IrClCOD) 5 mg exchanged	Saturated strontium	1 gm keisulghur was wetted with 100 μl portion of
	with TPPTS 100 mg.	chloride in 2 ml	solution A and evaporated under vaccume 10 mm Hg
	Poly acrylic acid sodium salt	water	with simultanious tumbling remining solution A was
	100 mg		added in 100 µl fractions and solid was isolated
	In 2 ml water		(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
			filtered to yield pale yellow colored solid powder.
125.	(IrClCOD) 5 mg exchanged	Saturated strontium	1 gm bentonite was wetted with 100 μl portion of
	with TPPTS 100 mg.	chloride in 2 ml	solution A and evaporated under vaccume 10 mm Hg
	Poly acrylic acid sodium salt	water	with simultanious tumbling remining solution A was
	100 mg		added in 100 µl fractions and solid was isolated
	In 2 ml water		(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
	•		filtered to yield pale yellow colored solid powder.
126.	(RuCl ₂ COD) 5 mg exchanged	Saturated strontium	1 gm davisil was wetted with 100 μl portion of solution
	with diphenyl phosphino	chloride in 2 ml	A and evaporated under vaccume 10 mm Hg with
	ethane tetrasulfonate 100 mg.	water	simultanious tumbling remining solution A was added
	Poly acrylic acid sodium salt		in 100 µl fractions and solid was isolated (moisture
	100 mg		
•			content ~20 %) this powder was added to solution B in
	In 2 ml water		equal fractions over a period of 2 hours and suspension
		•	was further agitated for 10 hours filtered to yield pale
			yellow colored solid powder.
127.	.(RuCl ₂ COD) 5 mg exchanged	Saturated strontium	1 gm davisil was wetted with 100 μl portion of solution
	with diphenyl phosphino	chloride in 2 ml	A and evaporated under vaccume 10 mm Hg with
	ethane tetrasulfonate 100 mg.	water	simultanious tumbling remining solution A was added
	Poly acrylic acid sodium salt	•	in 100 µl fractions and solid was isolated (moisture
	100 mg		content ~20 %) this powder was added to solution B in
	, ,		
	In 2 ml water		equal fractions over a period of 2 hours and suspension
	_		equal fractions over a period of 2 hours and suspension was further agitated for 10 hours filtered to yield pale
	_		•

	(P. 01 00P) 5	5001	1 gm davisil was wetted with 100 μl portion of solution
128.	(RuCl ₂ COD) 5 mg exchanged	500 mg calcium	A and evaporated under vaccume 10 mm Hg with
	with diphenyl phosphino	chloride in 2 ml	
	ethane tetrasulfonate 100 mg.	water	simultanious tumbling remining solution A was added
	Poly acrylic acid sodium salt		in 100 µl fractions and solid was isolated (moisture
	100 mg	4	content ~20 %) this powder was added to solution B in
	In 2 ml water		equal fractions over a period of 2 hours and suspension
			was further agitated for 10 hours filtered to yield light
	·	,	brown- yellow colored solid powder.
129.	Rh(COD)PF S,S chiraphos	Saturated strontium	1 gm davisil was wetted with 100 μl portion of solution
	tetrasulfonate 25 mg	chloride solution 2	A and evaporated under vaccume 10 mm Hg with
	S,S chiraphos tetrasulfonate	ml	simultanious tumbling remining solution A was added
	25 mg	·	in 100 µl fractions and solid was isolated (moisture
	Sodium alginate 100 mg		content ~20 %) this powder was added to solution B in
	dissolved in 2 ml water		equal fractions over a period of 2 hours and suspension
			was further agitated for 10 hours filtered to yield pale
			yellow colored solid powder.
	Rh(COD)PF S,S chiraphos	Saturated barium	1 gm davisil was wetted with 100 μl portion of solution
130.		nitrate solution 2 ml	A and evaporated under vaccume 10 mm Hg with
•	tetrasulfonate 25 mg S,S chiraphos tetrasulfonate	muac soluton 2 m	simultanious tumbling remining solution A was added
			in 100 µl fractions and solid was isolated (moisture
	25 mg		content ~20 %) this powder was added to solution B in
	Sodium alginate 100 mg		•
	dissolved in 2 ml water		equal fractions over a period of 2 hours and suspension
			was further agitated for 10 hours filtered to yield pale
			yellow colored solid powder.
131.	Rh(COD)PF ₆ / S,S chiraphos	Saturated barium	1 gm alumina was wetted with 100 μl portion of
	tetrasulfonate 25 mg	nitrate solution 2 ml	solution A and evaporated under vaccume 10 mm Hg
	S,S chiraphos tetrasulfonate		with simultanious tumbling remining solution A was
	25 mg		added in 100 µl fractions and solid was isolated
	Sodium alginate 100 mg		(moisture content ~20 %) this powder was added to
	dissolved in 2 ml water	· .	solution B in equal fractions over a period of 2 hours
	,		and suspension was further agitated for 10 hours
			filtered to yield pale yellow colored solid powder.
132.	Rh(COD)PF ₆ / S,S chiraphos	Saturated barium	1 gm titania was wetted with 100 μl portion of solution
	tetrasulfonate 25 mg	nitrate solution 2 ml	A and evaporated under vaccume 10 mm Hg with
	S,S chiraphos tetrasulfonate		simultanious tumbling remining solution A was added
	25 mg		in 100 µl fractions and solid was isolated (moisture
	Sodium alginate 100 mg		content ~20 %) this powder was added to solution B in
	dissolved in 2 ml water		equal fractions over a period of 2 hours and suspension
			was further agitated for 10 hours filtered to yield pale
,		·	yellow colored solid powder.
133.	HRhCO(TPATS) ₃	500 mg Calcium	1 gm titania was wetted with 100 μl portion of solution
133.		chloride solution in	A and evaporated under vaccume 10 mm Hg with
•	10 mg 100 mg TPATS	water 5 ml	simultanious tumbling remining solution A was added
		, , , , , , , , , , , , , , , , , , ,	·
	carboxy methyl cellulose		in 100 µl fractions and solid was isolated (moisture
	sodium 100 mg in 1 ml water		content ~20 %) this powder was added to solution B in
			equal fractions over a period of 2 hours and suspension
	-		was further agitated for 10 hours filtered to yield pale
	1		yellow-green colored solid powder.

134.	HRhCO(TPATS)3	Strontium chloride	1 gm alumina was wetted with 100 μl portion of
	10 mg	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	100 mg TPATS	water 5 ml	with simultanious tumbling remining solution A was
	carboxy methyl cellulose		added in 100 µl fractions and solid was isolated
	sodium 100 mg in 1 ml water		(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
	·		and suspension was further agitated for 10 hours
	i i		filtered to yield pale yellow-green colored solid
	·		powder.
135.	HRhCO(TPATS) ₃	Barium nitrate	1 gm bentonite was wetted with 100 μl portion of
100.	10 mg	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	100 mg TPATS	water 5 ml	with simultanious tumbling remining solution A was
	carboxy methyl cellulose	, water 5 km	added in 100 µl fractions and solid was isolated
	· · · · · · · · · · · · · · · · · · ·		· ·
	sodium 100 mg in 1 ml water		(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
		•	filtered to yield pale yellow-green colored solid
			powder.
136.	HRhCO(TPATS) ₃	Strontium chloride	1 gm titania was wetted with 100 μl portion of solution
	10 mg	saturated solution in	A and evaporated under vaccume 10 mm Hg with
	100 mg TPATS	water 5 ml	simultanious tumbling remining solution A was added
	carboxy methyl cellulose		in 100 µl fractions and solid was isolated (moisture
	sodium 100 mg in 1 ml water		content ~20 %) this powder was added to solution B in
			equal fractions over a period of 2 hours and suspension
			was further agitated for 10 hours filtered to yield pale
		,	yellow-green colored solid powder.
137.	HRbCO(TPATS)3	Strontium chloride	1 gm davisil was wetted with 100 μl portion of solution
	10 mg	saturated solution in	A and evaporated under vaccume 10 mm Hg with
	. 100 mg TPATS	water 5 ml	simultanious tumbling remining solution A was added
	carboxy methyl cellulose		in 100 µl fractions and solid was isolated (moisture
	sodium 100 mg in 1 ml water		content ~20 %) this powder was added to solution B in
			equal fractions over a period of 2 hours and suspension
·			was further agitated for 10 hours filtered to yield pale
			yellow-green colored solid powder.
138.	HRhCO(BISBIS) 50 mg	Saturated barium	2 gm davisil was wetted with 100 μl portion of solution
	BISBIS 200 mg	nitrate solution is 5	A and evaporated under vaccume 10 mm Hg with
	200 mg sodium sulfate	ml water	simultanious tumbling remining solution A was added
	dissolved in 2 ml water	. ,	in 100 µl fractions and solid was isolated (moisture
	dissolved in 2 in water		•
			content -20 %) this powder was added to solution B in
		i	equal fractions over a period of 2 hours and suspension
			was further agitated for 10 hours filtered to yield pale
		<u></u>	yellow colored solid powder.
139.	HRhCO(BISBIS) 50 mg	1 g calcium chloride	2 gm davisil was wetted with 100 μl portion of solution
	BISBIS 200 mg	solution In 5 ml	A and evaporated under vaccume 10 mm Hg with
	200 mg polyvinyl sulfonic	water	simultanious tumbling remining solution A was added
	acid		in 100 µl fractions and solid was isolated (moisture
	dissolved in 2 ml water		content -20 %) this powder was added to solution B in
			equal fractions over a period of 2 hours and suspension
			was further agitated for 10 hours filtered to yield pale
			yellow colored solid powder.

140.	HRhCO(BISBIS) 50 mg	Saturated barium	2 gm titania was wetted with 100 μl portion of solution
	BISBIS 200 mg	nitrate solution is 5	A and evaporated under vaccume 10 mm Hg with
	200 mg polyacrylic acid	ml water	simultanious tumbling remining solution A was added
	sodium salt		in 100 µl fractions and solid was isolated (moisture
	dissolved in 2 ml water		content ~20 %) this powder was added to solution B in
			equal fractions over a period of 2 hours and suspension
			was further agitated for 10 hours filtered to yield pale
		•	yellow colored solid powder.
141.	HRhCO(BISBIS) 50 mg	Saturated strontium	2 gm alumina was wetted with 100 μl portion of
	BISBIS 200 mg	chloride solution is 5	solution A and evaporated under vaccume 10 mm Hg
	200 mg polyvinyl sulfonic	ml water	with simultanious tumbling remining solution A was
	acid		added in 100 µl fractions and solid was isolated
	dissolved in 2 ml water	,	(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
			filtered to yield pale yellow colored solid powder.
142	HRhCO(BISBIS) 50 mg	Saturated barium	2 gm bentonite was wetted with 100 µl portion of
142.		nitrate solution is 5	solution A and evaporated under vaccume 10 mm Hg
	BISBIS 200 mg 200 mg polyvinyl sulfonic	ml water	with simultanious tumbling remining solution A was
		im water	added in 100 µl fractions and solid was isolated
	dissolved in 2 ml water		•
,	dissolved in 2 mi water		(moisture content ~20 %) this powder was added to
	•		solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
	<u> </u>		filtered to yield pale yellow colored solid powder.
143.	HRhCO(BISBIS) 50 mg	Saturated barium	2 gm davisil was wetted with 100 μl portion of solution
	BISBIS 200 mg	nitrate solution is 5	A and evaporated under vaccume 10 mm Hg with
	200 mg polyvinyl sulfonic	ml water	simultanious tumbling remining solution A was added
	acid		in 100 µl fractions and solid was isolated (moisture
:	dissolved in 2 ml water		content ~20 %) this powder was added to solution B in
			equal fractions over a period of 2 hours and suspension
		•	was further agitated for 10 hours filtered to yield pale
			yellow-green colored solid powder.
144.	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm davisil was wetted with 100 μl portion of solution
	TPPTS 100 mg	barium nitrate 5 ml	A and evaporated under vaccume 10 mm Hg with
	100 mg sodium alginate	į	simultanious tumbling remining solution A was added
	Dissolved in 2 ml water and		in 100 µl fractions and solid was isolated (moisture
	0.5 ml butane diol		content ~20 %) this powder was added to solution B in
i			equal fractions over a period of 2 hours and suspension
			was further agitated for 10 hours filtered to yield pale
			yellow-green colored solid powder.
145.	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm alumina was wetted with 100 μl portion of
	TPPTS 100 mg	barium nitrate 5 ml	solution A and evaporated under vaccume 10 mm Hg
	100 mg oxalic acid sodium		with simultanious tumbling remining solution A was
	salt.		added in 100 µl fractions and solid was isolated
	Dissolved in 2 ml water and		(moisture content ~20 %) this powder was added to
	0.5 ml butane diol		solution B in equal fractions over a period of 2 hours
	C.J III CAMIN MOI		
			and suspension was further agitated for 10 hours
			dilegand to unight male trailour necon animond animal
			filtered to yield pale yellow-green colored solid powder.

146.	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm davisil was wetted with 100 µl portion of solution
•	TPPTS 100 mg	strontium chloride 5	A and evaporated under vaccume 10 mm Hg with
	100 mg citric acid	ml	simultanious tumbling remining solution A was added
	Dissolved in 2 ml water and		in 100 µl fractions and solid was isolated (moisture
-	0.5 ml ethylene glycol		content ~20 %) this powder was added to solution B in
			equal fractions over a period of 2 hours and suspension
			was further agitated for 10 hours filtered to yield pale
			yellow-green colored solid powder.
147.	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm davisil was wetted with 100 μl portion of solution
	TPPTS 100 mg	barium nitrate 5 ml	A and evaporated under vaccume 10 mm Hg with
	100 mg polyacrylic acid		simultanious tumbling remining solution A was added
	sodium salt.		in 100 µl fractions and solid was isolated (moisture
	Dissolved in 2 ml water and	·	content ~20 %) this powder was added to solution B in
	0.5 ml butane diol		equal fractions over a period of 2 hours and suspension
		•	was further agitated for 10 hours filtered to yield pale
			yellow-green colored solid powder.
148.	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm shreded asbestos roap was wetted with 100 μl
	TPPTS 300 mg	barium nitrate 5 ml	portion of solution A and evaporated under vaccume 10
	Dissolved in 2 ml water	•	mm Hg with simultanious tumbling remining solution
	, -		A was added in 100 µl fractions and solid was isolated
	· · ·		(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
			filtered to yield pale yellow-green colored solid
			powder.
149.	Cobalt N, N'ethylene bis	Saturated barium	2 gm davisil was wetted with 100 μl portion of solution
	(salicyldiamine) 5-sulfonato	nitrate solution in	A and evaporated under vaccume 10 mm Hg with
	sodium 100 mg.	water 5ml	simultanious tumbling remining solution A was added
	Sodium phosphate. 500 mg.		in 100 µl fractions and solid was isolated (moisture
	In 5 ml water		content ~20 %) this powder was added to solution B in
			equal fractions over a period of 2 hours and suspension
• •			was further agitated for 10 hours filtered to yield pale
			brown colored solid powder.
150.	Cobalt N, N'ethylene bis	Saturated barium	2 gm alumina was wetted with 100 μl portion of
	(salicyldiamine) 5-sulfonato	nitrate solution in	solution A and evaporated under vaccume 10 mm Hg
:	sodium 100 mg.	water 5ml	with simultanious tumbling remining solution A was
	Sodium silicate 500 mg.		added in 100 µl fractions and solid was isolated
	In 5 ml water		(moisture content -20 %) this powder was added to
	·		solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
			filtered to yield pale brown colored solid powder.
151.	Cobalt N, N'ethylene bis	Saturated barium	2 gm titania was wetted with 100 μl portion of solution
÷	(salicyldiamine) 5-sulfonato	nitrate solution in	A and evaporated under vaccume 10 mm Hg with
	sodium 100 mg.	water 5ml	simultanious tumbling remining solution A was added
	Polyvinyl sulfonate sodium.		in 100 µl fractions and solid was isolated (moisture
	500 mg.		content ~20 %) this powder was added to solution B in
	In 5 ml water		equal fractions over a period of 2 hours and suspension
•			was further agitated for 10 hours filtered to yield pale
			brown colored solid powder.
			otown colored sond powder.

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152.	Cobalt N, N'ethylene bis	Saturated barium	2 gm zirconia was wetted with 100 µl portion of
	(salicyldiamine) 5-sulfonato	nitrate solution in	solution A and evaporated under vaccume 10 mm Hg
	sodium 100 mg.	water 5ml	with simultanious tumbling remining solution A was
	Polyvinyl sulfonate sodium.		added in 100 µl fractions and solid was isolated
	500 mg.		(moisture content ~20 %) this powder was added to
•	In 5 ml water		solution B in equal fractions over a period of 2 hours
·			and suspension was further agitated for 10 hours
			filtered to yield pale brown colored solid powder.
153.	Cobalt N, N'ethylene bis	2g calcium chloride	2 gm shreded asbestos roap was wetted with 100 μl
	(salicyldiamine) 5-sulfonato	solution in water 5ml	portion of solution A and evaporated under vaccume 10
	sodium 100 mg.		mm Hg with simultanious tumbling remining solution
	Polyvinyl sulfonate sodium.		A was added in 100 µl fractions and solid was isolated
	500 mg.		(moisture content ~20 %) this powder was added to
	In 5 ml water	·	solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
			filtered to yield gray colored solid powder.
154.	Cobalt (II), 4, 4', 4'',4'''-	Saturated strontium	2 gm shreded asbestos roap was wetted with 100 μl
134.	tetrasulfopthalocynine. 500	chloride in 5 ml	portion of solution A and evaporated under vaccume 10
		water	mm Hg with simultanious tumbling remining solution
	mg And 500 mg sodium sodium		A was added in 100 µl fractions and solid was isolated
		-	· ·
	poly vinyl sulfonate in 5 ml		(moisture content ~20 %) this powder was added to
	water		solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
			filtered to yield blue-gray colored solid powder.
155.	Cobalt (II), 4, 4', 4",4"'-	Saturated barium	2 gm kesilghur was wetted with 100 μl portion of
,	tetrasulfopthalocynine. 500	nitrate in 5 ml water	solution A and evaporated under vaccume 10 mm Hg
	mg		with simultanious tumbling remining solution A was
	And 500 mg sodium		added in 100 µl fractions and solid was isolated
	phosphate in 5 ml water		(moisture content ~20 %) this powder was added to
		·	solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
			filtered to yield pale blue colored solid powder.
156.	Cobalt (II), 4, 4', 4'',4'''-	Saturated strontium	2 gm kesilghur was wetted with 100 μl portion of
	tetrasulfopthalocynine. 500	chloride in 5 ml	solution A and evaporated under vaccume 10 mm Hg
	mg	water	with simultanious tumbling remining solution A was
	And 500 mg sodium		added in 100 µl fractions and solid was isolated
	phosphate in 5 ml water		(moisture content ~20 %) this powder was added to
`			solution B in equal fractions over a period of 2 hours
·	•		and suspension was further agitated for 10 hours
			filtered to yield pale blue colored solid powder.
157.		<u> </u>	
	Cobalt (II), 4, 4', 4'',4'''-	500mg. CaCl ₂ in 5 ml	2 gm kesilghur was wetted with 100 µl portion of
	tetrasulfopthalocynine. 500	water	solution A and evaporated under vaccume 10 mm Hg
	mg	•	with simultanious tumbling remining solution A was
• •	And 500 mg sodium		added in 100 µl fractions and solid was isolated
	phosphate in 5 ml water		(moisture content ~20 %) this powder was added to
	hroshuum iii 2 iiii mami		·
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
			filtered to yield pale blue colored solid powder.

	<u> </u>		
158.	Copper (II), 4, 4', 4",4"'-	500mg. CaCl₂ in 5 ml	2 gm kesilghur was wetted with 100 μl portion of
	tetrasulfopthalocynine. 500	water	solution A and evaporated under vaccume 10 mm Hg
	mg		with simultanious tumbling remining solution A was
	And 500 mg sodium sulfate in		added in 100 µl fractions and solid was isolated
	5 ml water		(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
			filtered to yield pale blue colored solid powder.
159.	Copper (II), 4, 4', 4'',4'''-	Saturated strontium	2 gm kesilghur was wetted with 100 μl portion of
•	tetrasulfopthalocynine. 500	chloride in 5 ml	solution A and evaporated under vaccume 10 mm Hg
	mg	water	with simultanious tumbling remining solution A was
	And 500 mg sodium silicate		added in 100 µl fractions and solid was isolated
	in 5 ml water		(moisture content ~20 %) this powder was added to
	'		solution B in equal fractions over a period of 2 hours
	,		and suspension was further agitated for 10 hours
	·		filtered to yield pale blue colored solid powder.
160.	Copper (II), 4, 4', 4'',4'''-	Saturated barium	2 gm kesilghur was wetted with 100 μl portion of
	tetrasulfopthalocynine. 500	nitrate in 5 ml water	solution A and evaporated under vaccume 10 mm Hg
	mg	·	with simultanious tumbling remining solution A was
	And 500 mg sodium silicate		added in 100 µl fractions and solid was isolated
	in 5 ml water		(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
	· · ·		and suspension was further agitated for 10 hours
		·	filtered to yield pale blue colored solid powder.
161.	Copper (II), 4, 4', 4", 4"'-	Saturated barium	2 gm bentonite was wetted with 100 µl portion of
	tetrasulfopthalocynine. 500	nitrate in 5 ml water	solution A and evaporated under vaccume 10 mm Hg
•	mg		with simultanious tumbling remining solution A was
	And 500 mg sodium silicate	·	added in 100 µl fractions and solid was isolated
	in 5 ml water		(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
			filtered to yield pale blue colored solid powder.
162.	Copper (II), 4, 4', 4'',4'''-	Saturated strontiun	2 gm bentonite was wetted with 100 µl portion of
	tetrasulfopthalocynine. 500	chloride in 5 ml	solution A and evaporated under vaccume 10 mm Hg
	mg	water	with simultanious tumbling remining solution A was
	And 500 mg sodium silicate	,	added in 100 µl fractions and solid was isolated
:	in 5 ml water		(moisture content ~20 %) this powder was added to
		•	solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
	,		filtered to yield pale blue colored solid powder.
163.			
	Manganese(II), 4, 4', 4'',4'''-	Saturated strontiun	2 gm davisil was wetted with 100 μl portion of solution
	tetrasulfopthalocynine. 500	chloride in 5 ml	A and evaporated under vaccume 10 mm Hg with
	mg	water	simultanious tumbling remining solution A was added
	And 500 mg sodium silicate		in 100 µl fractions and solid was isolated (moisture
	in 5 ml water	· 	content ~20 %) this powder was added to solution B in
			equal fractions over a period of 2 hours and suspension
			was further agitated for 10 hours filtered to yield pale
			blue colored solid powder.

164.	Manganese(II), 4, 4', 4",4"'-	Saturated barium	2 gm davisil was wetted with 100 µl portion of solution
104.	tetrasulfopthalocynine. 500	nitrate in 5 ml water	A and evaporated under vaccume 10 mm Hg with
•	mg		simultanious tumbling remining solution A was added
	And 500 mg sodium silicate		in 100 µl fractions and solid was isolated (moisture
-	in 5 ml water		content ~20 %) this powder was added to solution B in
	in 5 ini wates		equal fractions over a period of 2 hours and suspension
			was further agitated for 10 hours filtered to yield pale
	277 4 41 411 411	Samuel Links	blue colored solid powder.
165.	Manganese(II), 4, 4', 4'',4'''-	Saturated barium	2 gm alumina was wetted with 100 µl portion of
	tetrasulfopthalocynine. 500	nitrate in 5 ml water	solution A and evaporated under vaccume 10 mm Hg
,	mg		with simultanious tumbling remining solution A was
	And 500 mg sodium silicate		added in 100 µl fractions and solid was isolated
	in 5 ml water		(moisture content ~20 %) this powder was added to
			solution B in equal fractions over a period of 2 hours
			and suspension was further agitated for 10 hours
			filtered to yield pale blue colored solid powder.
166.	Manganese(II), 4, 4', 4",4"'-	Saturated barium	2 gm alumina was wetted with 100 μl portion of
	tetrasulfopthalocynine. 500	nitrate in 5 ml water	solution A and evaporated under vaccume 10 mm Hg
	mg		with simultaneous tumbling remaining solution A was
	And 500 mg sodium		added in 100 µl fractions and solid was isolated
	polyvinyl sulfonate in 5 ml	÷	(moisture content ~20 %) this powder was added to
	water		solution B in equal fractions over a period of 2 hours
	·		and suspension was further agitated for 10 hours
	ľ		filtered to yield pale blue colored solid powder.
167.	Iron (III), 4, 4', 4'',4'''-	Saturated strontium	2 gm davisil was wetted with 100 μl portion of solution
	tetrasulfopthalocynine oxygen	chloride in 5 ml	A and evaporated under vaccume 10 mm Hg with
	adduct. 500 mg	water	simultanious tumbling remining solution A was added
	And 500 mg sodium sulfate in		in 100 µl fractions and solid was isolated (moisture
	5 ml water		content ~20 %) this powder was added to solution B in
•		·	equal fractions over a period of 2 hours and suspension
			was further agitated for 10 hours filtered to yield pale
			blue colored solid powder.
168.	Iron (III), 4, 4', 4'',4'''-	Saturated barium	2 gm davisil was wetted with 100 μl portion of solution
	tetrasulfopthalocynine oxygen	nitrate in water 5 ml	A and evaporated under vaccume 10 mm Hg with
	adduct. 500 mg		simultanious tumbling remining solution A was added
	And 500 mg sodium sulfate in		in 100 µl fractions and solid was isolated (moisture
	5 ml water		content ~20 %) this powder was added to solution B in
:			equal fractions over a period of 2 hours and suspension
			was further agitated for 10 hours filtered to yield pale
		,	blue colored solid powder.

Examples 169 to 252

Preparation of catalytic formulation by deposition precipitation with simultaneous removal of water.

The following examples illustrate one of the procedures for the preparation of the catalytic formulation of the invention in accordance with the method of formulation known as co precipitation near the surface of the solid support.

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The general procedure for the preparation of heterogeneous catalytic formulation is described herein as making of a solution of anionically charged catalytic entity, catalytically inert anionic additive (termed as solution A) and solution of group II A metal ions (termed as solution B). The specified amount of support pretreated as described in earlier is impregnated with solution A by wetting solids with solution followed by evaporation to obtain dry solid support bearing anionic components of solution A. This solid powder was suspended in water immiscible solvent or a solvent that forms azeotrope with solvent component of solution B. the suspension was agitated and temperature was raised such that solvent starts distilling. Under this condition solution B was slowly pumped in. simultaneously solvent in which solids are suspended is also pumped in with rate similar to that of distillation. Once all solution B was added suspension was stirred for specified period of time. Resulting suspension is further agitated for specified time. Suspension was centrifuged and solids were repeatedly washed with water and dried in vacuum. Dry powder was stored under argon in gas tight vessel. These solid catalytic formulations can be used for appropriate reactions depending upon catalytically active entity incorporated in it.

Note1: solution A is prepared by dissolving anionic components including anionic complex and additives to make homogeneous solution in degassed solvents. And purging argon also degasses resulting solution.

- Note 2: solution B is prepared by dissolving dissolving group IIA metal salts. Solution was degassed prior to use
- Note 3: the impregnation of solution on solid support is carried out by wetting solids with solution A and evaporating in vacuum at 50 °C unless stated
- Note 4: addition of impregnated solids with components of A to solution B is carried out at ambient temperature unless stated.
 - Note 5: the impregnation of solution A may be bypassed instead following procedure may be employed. Support is suspended in solvent to which solution is pumped in with simultaneous removal of solvent component of solution A. solvent is also pumped in such a rate that liquid volume of the container remain same. After this solution B addition aging and solid isolation is carried out as described earlier.

T	Colution A	Solution B	Procedure
Example	Solution A	Solution B Saturated barium	2 gm Davisil was wetted with 100 µl portion of
169 ·	HRhCO(TPPTS)3, 50 mg,	nitrate in water 2 ml	solution A and evaporated under vaccume 10 mm Hg
	TPPTS 200 mg. Dissolved in water 2 ml	uittate in water 2 im	with simultanious tumbling remining solution A was
	Dissolved in water 2 mi		added in 100 µl fractions and solid was isolated this
	•		powder was added to benzene 25 ml in apparatus
			described in fig.relux was started and solution B was
	•		added in equal fractions over a period of 2 hours while
			simultanious removal of azeotropic waer and
*	,		suspension was further agitated for 10 hours to yild
			pale yellow colored solid powder.
170	HRhCO(TPPTS)3, 50 mg,	Saturated strontium	2 gm Davisil was wetted with 100 μl portion of
!	TPPTS 200 mg.	chloride in water 2ml	solution A and evaporated under vaccume 10 mm Hg
	Dissolved in water 2 ml	ļ	with simultanious tumbling remining solution A was
			added in 100 µl fractions and solid was isolated this
	•		powder was added to benzene 25 ml in apparatus
			described in fig.relux was started and solution B was
			added in equal fractions over a period of 2 hours while
•		•	simultanious removal of azeotropic waer and
			suspension was further agitated for 10 hours to yild
*		500 - 61-	pale yellow colored solid powder.
171	HRhCO (TPPTS)3, 50 mg,	500 mg of calcium	2 gm Davisil was wetted with 100 µl portion of
	TPPTS 200 mg.	chloride in 2 ml	solution A and evaporated under vaccume 10 mm Hg
	Dissolved in water 2 ml	water	with simultanious tumbling remining solution A was
	:		added in 100 µl fractions and solid was isolated this
	•		powder was added to benzene 25 ml in apparatus
	·		described in fig.relux was started and solution B was
			added in equal fractions over a period of 2 hours while
	•		simultanious removal of azeotropic waer and suspension was further agitated for 10 hours to yild
	,		pale yellow colored solid powder.
150	LIDLOO (TDPTC)2 60 mg	Barium nitrate	2 gm γ-alumina was wetted with 100 μl portion of
172	HRhCO (TPPTS)3, 50 mg,	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	TPPTS 200 mg.	water	with simultanious tumbling remining solution A was
	Dissolved in water 2 ml	Watci	added in 100 µl fractions and solid was isolated this
			powder was added to benzene 25 ml in apparatus
:			described in fig.relux was started and solution B was
		•	added in equal fractions over a period of 2 hours while
			simultanious removal of azeotropic waer and
•			suspension was further agitated for 10 hours to yild
			pale yellow colored solid powder.
172	UBPCO(LDDLe)3 20 ma	Strontium chloride	2 gm γ-alumina was wetted with 100 μl portion of
173	HRhCO(TPPTS)3, 50 mg,	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	TPPTS 200 mg. Dissolved in water 2 ml	water 2ml	with simultanious tumbling remining solution A was
**	Dissured in water 2 iii		added in 100 µl fractions and solid was isolated this
			powder was added to benzene 25 ml in apparatus
			described in fig.relux was started and solution B was
		ŀ	added in equal fractions over a period of 2 hours while
			simultanious removal of azeotropic waer and
			suspension was further agitated for 10 hours to yild
			pale yellow colored solid powder.
I	l .	I	Pare Jenew colored sould powder.

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174.	HRhCO(TPPTS)3, 50 mg,	Calcium chloride 500	2 gm γ-alumina was wetted with 100 μl portion of
·	TPPTS 200 mg.	mg solution in 2 ml	solution A and evaporated under vaccume 10 mm Hg
	Dissolved in water 2 ml	water	with simultanious tumbling remining solution A was
	1	1	added in 100 µl fractions and solid was isolated this
			powder was added to benzene 25 ml in apparatus
			described in fig relux was started and solution B was
			added in equal fractions over a period of 2 hours while
	·		simultanious removal of azeotropic waer and
	1		suspension was further agitated for 10 hours to yild
			pale yellow colored solid powder.
	11D1 00 (TDDTTTC) 0 50	<u> </u>	
175.	HRhCO(TPPTS)3, 50 mg,	Barium nitrate	2 gm bentonite was wetted with 100 μl portion of
	TPPTS 200 mg.	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	Dissolved in water 2 ml	water 2 ml	with simultanious tumbling remining solution A was
			added in 100 µl fractions and solid was isolated this
			powder was added to benzene 25 ml in apparatus
	•		described in fig.relux was started and solution B was
,			added in equal fractions over a period of 2 hours while
			simultanious removal of azeotropic waer and
			suspension was further agitated for 10 hours to yild
			pale yellow colored solid powder.
176.	HRhCO(TPPTS)3, 50 mg,	Strontium chloride	2 gm bentonite was wetted with 100 μl portion of
	TPPTS 200 mg.	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
•	Dissolved in water 2 ml	water	with simultanious tumbling remining solution A was
		,	added in 100 µl fractions and solid was isolated this
			powder was added to benzene 25 ml in apparatus
		,	•
			described in fig.relux was started and solution B was
			added in equal fractions over a period of 2 hours while
			simultanious removal of azeotropic waer and
ŀ			suspension was further agitated for 10 hours to yild
			pale yellow colored solid powder.
177.	HRhCO (TPPTS)3, 50 mg,	Calcium chloride 500	2 gm bentonite was wetted with 100 μl portion of
	TPPTS 200 mg.	mg solution in 2 ml	solution A and evaporated under vaccume 10 mm Hg
	Dissolved in water 2 ml	water	with simultanious tumbling remining solution A was
Į			added in 100 µl fractions and solid was isolated this
		:	powder was added to benzene 25 ml in apparatus
	,		described in fig.relux was started and solution B was
		·	added in equal fractions over a period of 2 hours while
			simultanious removal of azeotropic waer and
			suspension was further agitated for 10 hours to yild
		•	pale yellow colored solid powder.
178.	HRhCO (TPPTS)3, 50 mg,	Barium nitrate	2 gm charcoal was wetted with 100 μl portion of
170.	TPPTS 200 mg.	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
·	Dissolved in water 2 ml	water	·
	Pissoived III Water 2 IIII		with simultanious tumbling remining solution A was
·			added in 100 µl fractions and solid was isolated this
		,	powder was added to benzene 25 ml in apparatus
	·		described in fig.relux was started and solution B was
			added in equal fractions over a period of 2 hours while
,			simultanious removal of azeotropic waer and
·			suspension was further agitated for 10 hours to yild
			black colored solid powder.
			

179.	HRhCO (TPPTS)3, 50 mg,	Strontium chloride	2 gm charcoal was wetted with 100 μl portion of
179.	TPPTS 200 mg.	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
•		water	with simultanious tumbling remining solution A was
	Dissolved in water 2 ml	watti	added in 100 µl fractions and solid was isolated this
		;	l · · · · · · · · · · · · · · · · · · ·
* *			powder was added to benzene 25 ml in apparatus
			described in fig.relux was started and solution B was
		•	added in equal fractions over a period of 2 hours while
			simultanious removal of azeotropic waer and
	•		suspension was further agitated for 10 hours to yild
			black colored solid powder.
180.	HRhCO(TPPTS)3, 50 mg,	Calcium chloride 500	2 gm charcoal was wetted with 100 µl portion of
	TPPTS 200 mg.	mg solution in 2 ml	solution A and evaporated under vaccume 10 mm Hg
	Dissolved in 2 ml water	water	with simultanious tumbling remining solution A was
			added in 100 µl fractions and solid was isolated this
	,		powder was added to benzene 25 ml in apparatus
			described in fig.relux was started and solution B was
		,	added in equal fractions over a period of 2 hours while
			simultanious removal of azeotropic waer and
			suspension was further agitated for 10 hours to yild
			black colored solid powder.
181.	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	2 gm davisil was wetted with 100 µl portion of solution
-	TPPTS 200 mg	saturated solution in	A and evaporated under vaccume 10 mm Hg with
	Dissolved in 2 ml water	2 ml water	simultanious tumbling remining solution A was added
	•		in 100 µl fractions and solid was isolated this powder
			was added to benzene 25 ml in apparatus described in
			fig.relux was started and solution B was added in equal
	•		fractions over a period of 2 hours while simultanious
			removal of azeotropic waer and suspension was further
			agitated for 10 hours filtured to yield light brown
			colored solid powder.
182.	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Strontium chloride	2 gm davisil was wetted with 100 μl portion of
	TPPTS 200 mg	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	Dissolved in 2 ml water.	2 ml water	with simultanious tumbling remining solution A was
			added in 100 µl fractions and solid was isolated this
	·		powder was added to benzene 25 ml in apparatus
			described in fig.relux was started and solution B was
			added in equal fractions over a period of 2 hours while
			simultanious removal of azeotropic waer and
			suspension was further agitated for 10 hours filtured to
•	•		yield light brown colored solid powder
183.	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	2 gm γ-alumina was wetted with 100 μl portion of
105.	TPPTS 200 mg Dissolved in	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	2 ml water	2 ml water	with simultanious tumbling remining solution A was
	. All Water	_ m	. =
·			added in 100 µl fractions and solid was isolated this
			powder was added to benzene 25 ml in apparatus
			described in fig.relux was started and solution B was
.	•		added in equal fractions over a period of 2 hours while
· 			simultanious removal of azeotropic waer and
•			suspension was further agitated for 10 hours filtered to
			yield light brown colored solid powder.

184.	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Strontium chloride	2 gm γ-alumina was wetted with 100 μl portion of
	TPPTS 200 mg Dissolved in	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	2 ml water	2 ml water	with simultanious tumbling remining solution A was
		·	added in 100 µl fractions and solid was isolated this
	•		powder was added to benzene 25 ml in apparatus
			described in fig.relux was started and solution B was
			added in equal fractions over a period of 2 hours while
			simultanious removal of azeotropic waer and
			suspension was further agitated for 10 hours filtered to
•	•	,	yield light brown colored solid powder.
185.	Ru(H)(Cl)(TPPTS) ₂ 50 mg	Strontium chloride	2 gm γ-alumina was wetted with 100 μl portion of
	TPPTS 200 mg	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	Sodium polyvinylsulfonate	2 ml water	with simultanious tumbling remining solution A was
	500 mg Dissolved in 2 ml		added in 100 µl fractions and solid was isolated this
	water		powder was added to benzene 25 ml in apparatus
			described in fig.relux was started and solution B was
		· .	added in equal fractions over a period of 2 hours while
			simultanious removal of azeotropic waer and
			suspension was further agitated for 10 hours filtered to
			yield light brown colored solid powder.
186.	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	2 gm γ-alumina was wetted with 100 μl portion of
2001	TPPTS 200 mg	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	Sodium polyvinylsulfonate	2 ml water	with simultanious tumbling remining solution A was
	500 mg Dissolved in 2 ml		added in 100 µl fractions and solid was isolated this
	water		powder was added to benzene 25 ml in apparatus
			described in fig.relux was started and solution B was
			added in equal fractions over a period of 2 hours while
			simultanious removal of azeotropic waer and
		•	suspension was further agitated for 10 hours filtered to
			yield light brown colored solid powder.
187.	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	2 gm titania was wetted with 100 μl portion of
1071	TPPTS 200 mg	saturated solution in	solution A and evaporated under vaccume 10 mm Hg
	Sodium polyvinylsulfonate	2 ml water	with simultanious tumbling remining solution A was
	500 mg Dissolved in 2 ml		added in 100 µl fractions and solid was isolated this
	water	,	powder was added to benzene 25 ml in apparatus
			described in fig.relux was started and solution B was
			added in equal fractions over a period of 2 hours while
			simultanious removal of azeotropic waer and
			suspension was further agitated for 10 hours filtered to
			yield light brown colored solid powder.
188.	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	2 gm zirconia 2 gm γ-alumina was wetted with 100 μl
100.	TPPTS 200 mg	saturated solution in	portion of solution A and evaporated under vaccume 10
	Sodium polyvinylsulfonate	2 ml water	mm Hg with simultanious tumbling remining solution
	500 mg Dissolved in 2 ml	_ <u></u>	
	-		A was added in 100 µl fractions and solid was isolated
·	water		this powder was added to benzene 25 ml in apparatus
			described in fig.relux was started and solution B was
			added in equal fractions over a period of 2 hours while
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			simultanious removal of azeotropic waer and
			simultanious removal of azeotropic waer and suspension was further agitated for 10 hours filtered to yield light brown colored solid powder.

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189.	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	2 gm activated charcoal was wetted with 100 µl portion
	TPPTS 200 mg	saturated solution in	of solution A and evaporated under vaccume 10 mm
	Sodium polyvinylsulfonate	2 ml water	Hg with simultanious tumbling remining solution A
	500 mg Dissolved in 2 ml	!	was added in 100 µl fractions and solid was isolated
	water		this powder was added to benzene 25 ml in apparatus
İ			described in fig.relux was started and solution B was
			added in equal fractions over a period of 2 hours while
			simultanious removal of azeotropic water and
		·	suspension was further agitated for 10 hours filtered to
			yield black colored solid powder.
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190.	PdCl ₂ (TPPTS) ₂ 10 mg	Barium nitrate	2 gm shreaded asbestos roap was suspended in benzene
	TPPTS 100 mg	saturated solution 5	25 ml in apparatus described in fig agitated. The
	Poly acrylic acid sodium salt	ml	temperature of the suspension was slowly raised such
	in 5 ml		that it gently refluxes to which was added 100 µl
		4	portion of solution A and solvent component was
			azeotropically removed, remining solution A was
			added in 100 µl fractions untill uniform suspension
			volume of suspension was maintained while
1			maintaining volume by pumping benzene and solution
			B was added in equal fractions over a period of 2 hours
			and simultanious removal of azeotropic water and
·			suspension was further continued. Formed suspension
•			was agitated.10 hours filtered to yield yellow gray
			colored solid powder.
191.	PdCl ₂ (TPPTS) ₂ 10 mg	Strontium chloride	2 gm shreaded asbestos roap was suspended in benzene
	TPPTS 100 mg	saturated solution 5	25 ml in apparatus described in fig agitated. The
	Poly acrylic acid sodium salt	ml	temperature of the suspension was slowly raised such
	in 5 ml		that it gently refluxes to which was added 100 µl
1			portion of solution A and solvent component was
			azeotropically removed, remining solution A was
			added in 100 µl fractions untill uniform suspension
			volume of suspension was maintained while
		•	maintaining volume by pumping benzene and solution
			B was added in equal fractions over a period of 2 hours
			and simultanious removal of azeotropic water and
			suspension was further continued. Formed suspension
,			was agitated for 10 hours filtered to yield yellow gray
			colored solid powder.
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192.	PdCl ₂ (TPPTS) ₂ 10 mg	500 mg	calcium	2 gm shreaded asbestos roap was suspended in
	TPPTS 100 mg	chloride ii	n 5 ml	benzene 25 ml in apparatus described in fig agitated.
	Poly acrylic acid sodium salt	water.		The temperature of the suspension was slowly raised
•	in 5 ml			such that it gently refluxes to which was added 100 μl
·				portion of solution A and solvent component was
				azeotropically removed, remining solution A was
				added in 100 µl fractions untill uniform suspension
		ľ		volume of suspension was maintained while
				maintaining volume by pumping benzene.and solution
Ì				B was added in equal fractions over a period of 2 hours
İ				
	•			and simultanious removal of azeotropic water and
				suspension was further continued. Formed suspension
				was agitated for 10 hours filtered to yield yellow gray
				colored solid powder.
193.	PdAc₂BYPYDS 25 mg	Barium	nitrate	2 gm davisil was suspended in benzene 25 ml in
Ì	BYPYDS 100 mg	saturated	solution	apparatus described in fig agitated. The temperature of
	Dissolved in 2 ml water	5ml		the suspension was slowly raised such that it gently
				refluxes to which was added 100 µl portion of solution
				A and solvent component was azeotropically removed,
٠	•			remining solution A was added in 100 μl fractions
				untill uniform suspension volume of suspension was
•				maintained while maintaining volume by pumping
				benzene and solution B was added in equal fractions
				over a period of 2 hours and simultanious removal of
	ι			azeotropic water and suspension was further continued.
				Formed suspension was agitated for 10 hours filtered to
				yield loght orange colored solid powder.
194.	PdAc ₂ BYPYDS 25 mg	Strontium	chloride	2 gm davisil was suspended in benzene 25 ml in
	BYPYDS 100 mg	saturated	solution	apparatus described in fig agitated. The temperature of
	Dissolved in 2 ml water	5ml		the suspension was slowly raised such that it gently
1				refluxes to which was added 100 µl portion of solution
	·			A and solvent component was azeotropically removed,
				remining solution A was added in 100 µl fractions
	,			untill uniform suspension volume of suspension was
				maintained while maintaining volume by pumping
				benzene and solution B was added in equal fractions
				over a period of 2 hours and simultanious removal of
				azeotropic water and suspension was further continued.
				Formed suspension was agitated for 10 hours filtered to
				yield loght orange colored solid powder.
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195.	PdAc ₂ BYPYDS 25 mg	500 mg calcium	2 gm davisil was suspended in benzene 25 ml in
	BYPYDS 100 mg	chloride in 5 ml	apparatus described in fig agitated. The temperature of
	Dissolved in 2 ml water	water	the suspension was slowly raised such that it gently
			refluxes to which was added 100 µl portion of solution
	· ·		A and solvent component was azeotropically removed,
			remining solution A was added in 100 µl fractions
		,	untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
			benzene and solution B was added in equal fractions
	·		over a period of 2 hours and simultanious removal of
	·		azeotropic water and suspension was further continued.
		,	Formed suspension was agitated for 10 hours filtered to
			yield loght orange colored solid powder.
196.	PdAc ₂ BYPYDS 25 mg	Barium nitrate	2 gm bentonite was suspended in benzene 25 ml in
	BYPYDS 100 mg	saturated solution	apparatus described in fig agitated. The temperature of
·	Dissolved in 2 ml water	5ml	the suspension was slowly raised such that it gently
			refluxes to which was added 100 µl portion of solution
		•	A and solvent component was azeotropically removed,
			remining solution A was added in 100 µl fractions
			untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
•			benzene and solution B was added in equal fractions
		•	over a period of 2 hours and simultanious removal of
			azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours filtered to
*			yield loght orange colored solid powder.
197.			
	PdAc₂ tri ortho tolyl	Barium nitrate	2 gm bentonite was suspended in benzene 25 ml in
	phosphine trisulfonated 25	saturated solution	apparatus described in fig agitated. The temperature of
	mg	5ml	the suspension was slowly raised such that it gently
	Tri ortho tolyl phosphine		refluxes to which was added 100 µl portion of solution
	trisulfonated 100 mg		A and solvent component was azeotropically removed,
	Dissolved in 2 ml water	·	remining solution A was added in 100 µl fractions
	·		untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
	·		benzene and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
,			
,	•		azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours filtered to
			yield pale yellow brown colored solid powder.
198.	PdAc ₂ tri ortho tolyl phosphine	Strontium chloride	2 gm bentonite was suspended in benzene 25 ml in apparatus
	trisulfonated 25 mg	saturated solution 5ml	described in fig agitated. The temperature of the suspension
	Tri ortho tolyl phosphine		was slowly raised such that it gently refluxes to which was
	trisulfonated 100 mg		added 100 µl portion of solution A and solvent component was
	Dissolved in 2 ml water		azeotropically removed, remining solution A was added in 100 · µl fractions untill uniform suspension volume of suspension
			was maintained while maintaining volume by pumping benzene
	,		and solution B was added in equal fractions over a period of 2
			hours and simultanious removal of azeotropic water and
			suspension was further continued. Formed suspension was
			agitated for .10 hours filtered to yield pale yellow brown
			colored solid powder.
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199.	PdAc ₂ trio tolyl phosphine trisulfonated 25 mg Tri ortho tolyl phosphine trisulfonated 100 mg Dissolved in 2 ml water	Barium nitrate saturated solution 5ml	2 gm alumina was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for 10 hours filtered to yield pale yellow brown colored solid powder.
200.	PdAc ₂ tri ortho tolyl phosphine trisulfonated 25 mg Tri ortho tolyl phosphine trisulfonated 100 mg Dissolved in 2 ml water	Barium nitrate saturated solution 5ml	2 gm charcoal was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for24 hours filtered to yield black colored solid powder.
201.	NiCl ₂ (TPPTS) ₂ 25 mg TPPTS 100 mg Sodium carboxy methyl cellulose 100 mg Dissolved in 2 ml	Saturated barium nitrate in 2 ml water	1 gm davisil was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 μl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 μl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for 10 hours filtered to yield pale solid powder of white color with blue tinge.

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202.	NiCl ₂ .(TPPTS) ₂ 25 mg TPPTS 100 mg Sodium carboxy methyl cellulose 100 mg Dissolved in 2 ml	Saturated barium nitrate in 2 ml water	1 gm alumina was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 μl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 μl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for for 10 hours
			filtered to yield solid powder of white color with blue tinge.
203	NiCl ₂ (TPPTS) ₂ 25 mg TPPTS 100 mg Sodium carboxy methyl cellulose 100 mg Dissolved in 2 ml	Saturated barium nitrate in 2 ml water	1 gm zirconia was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 μl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 μl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for 10 hours filtered to yield pale blue colored solid powder.

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204.	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated strontium	1 gm zirconia was suspended in benzene 25 ml in
	TPPTS 100 mg	chloride in 2 ml	apparatus described in fig agitated. The temperature of
j	Sodium carboxy methyl	water	the suspension was slowly raised such that it gently
1	cellulose 100 mg		refluxes to which was added 100 µl portion of solution
	Dissolved in 2 ml		A and solvent component was azeotropically removed,
			remining solution A was added in 100 μl fractions
	•		untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
			benzene and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
	·		azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours filtered to
	**	·	yield solid powder of white color with slight blue
	N'OL (TRIPTE) 25	Saturated strontium	tinge. 1 gm titania was suspended in benzene 25 ml in apparatus
205.	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated strontium chloride in 2 ml water	described in fig agitated. The temperature of the suspension
	TPPTS 100 mg Sodium carboxy methyl cellulose	Oldoride in 2 iin water	was slowly raised such that it gently refluxes to which was
	100 mg	·	added 100 µl portion of solution A and solvent component was
	Dissolved in 2 ml		azeotropically removed, remining solution A was added in 100
			μl fractions untill uniform suspension volume of suspension
			was maintained while maintaining volume by pumping benzene
			and solution B was added in equal fractions over a period of 2
			hours and simultanious removal of azeotropic water and
	·		suspension was further continued. Formed suspension was
	.		agitated for 10 hours filtered to yield solid powder of white
			color with blue tinge.
206.	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated strontium	1 gm asbestos was suspended in benzene 25 ml in apparatus
	TPPTS 100 mg	chloride in 2 ml water	described in fig agitated. The temperature of the suspension
•	Sodium carboxy methyl cellulose		was slowly raised such that it gently refluxes to which was
	100 mg		added 100 µl portion of solution A and solvent component was
	Dissolved in 2 ml		azeotropically removed, remining solution A was added in 100
			µl fractions untill uniform suspension volume of suspension
			was maintained while maintaining volume by pumping benzene
			and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and
			suspension was further continued. Formed suspension was
			agitated for 10 hours filtered to yield gray colored solid
			powder.
207.	(IrClCOD) 5 mg exchanged with	Saturated strontium	I gm davisil was suspended in benzene 25 ml in apparatus
	TPPTS 100 mg.	chloride in 2 ml water	described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was
	Poly acrylic acid sodium salt 100		added 100 µl portion of solution A and solvent component was
	In 2 ml water		azeotropically removed, remining solution A was added in 100
			ul fractions untill uniform suspension volume of suspension
			was maintained while maintaining volume by pumping benzene
1			and solution B was added in equal fractions over a period of 2
			hours and simultanious removal of azeotropic water and
			suspension was further continued. Formed suspension was
	,		agitated for 10 hours filtered to yield pale yellow colored solid
1			powder.
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	(I-C)COD) 5	Cotumbod	1 am baisulakur una suspandad in bangana 25 ml in
208.	(IrClCOD) 5 mg exchanged	Saturated strontium	1 gm keisulghur was suspended in benzene 25 ml in
	with TPPTS 100 mg.	chloride in 2 ml	apparatus described in fig agitated. The temperature of
•	Poly acrylic acid sodium salt	water	the suspension was slowly raised such that it gently
	100 mg		refluxes to which was added 100 µl portion of solution
	In 2 ml water		A and solvent component was azeotropically removed,
			remining solution A was added in 100 µl fractions
			untill uniform suspension volume of suspension was
<i>.</i>			maintained while maintaining volume by pumping
			benzene and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
			azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours filtered to
			yield pale yellow colored solid powder.
209.	(IrClCOD) 5 mg exchanged	Saturated strontium	1 gm bentonite was suspended in benzene 25 ml in
	with TPPTS 100 mg.	chloride in 2 ml	apparatus described in fig agitated. The temperature of
	Poly acrylic acid sodium salt	water	the suspension was slowly raised such that it gently
].	100 mg		refluxes to which was added 100 µl portion of solution
	In 2 ml water		A and solvent component was azeotropically removed,
	,		remining solution A was added in 100 µl fractions
	,		untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
			benzene and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
			azeotropic water and suspension was further continued.
			Formed suspension was agitated 10 hours filtered to
			yield pale yellow colored solid powder.
210.	(RuCl ₂ COD) 5 mg exchanged	Saturated strontium	1 gm davisil was suspended in benzene 25 ml in apparatus
,	with diphenyl phosphino ethane	chloride in 2 ml water	described in fig agitated. The temperature of the suspension
	tetrasulfonate 100 mg. Poly acrylic acid sodium salt 100		was slowly raised such that it gently refluxes to which was
	mg		added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100
	In 2 ml water		µl fractions untill uniform suspension volume of suspension
			was maintained while maintaining volume by pumping benzene
			and solution B was added in equal fractions over a period of 2
			hours and simultanious removal of azeotropic water and
			suspension was further continued. Formed suspension was
			agitated for 10 hours filtered to yield pale yellow-brown
	(P. Cl COP) 5	6	colored solid powder.
211.	(RuCl ₂ COD) 5 mg exchanged	Saturated strontium	I gm davisil was suspended in benzene 25 ml in
	with diphenyl phosphino	chloride in 2 ml	apparatus described in fig agitated. The temperature of
]	ethane tetrasulfonate 100 mg.	water	the suspension was slowly raised such that it gently
	Poly acrylic acid sodium salt		refluxes to which was added 100 µl portion of solution
	100 mg		A and solvent component was azeotropically removed,
	In 2 ml water		remining solution A was added in 100 μl fractions
			untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
	•		benzene and solution B was added in equal fractions
			l awar a mariad of 2 house and simultanious assessed of
			over a period of 2 hours and simultanious removal of
			azeotropic water and suspension was further continued.

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212.	(RuCl ₂ COD) 5 mg exchanged	500 mg calcium	1 gm davisil was suspended in benzene 25 ml in
	with diphenyl phosphino	chloride in 2 ml	apparatus described in fig agitated. The temperature of
	ethane tetrasulfonate 100 mg.	water	the suspension was slowly raised such that it gently
	Poly acrylic acid sodium salt		refluxes to which was added 100 µl portion of solution
	100 mg		A and solvent component was azeotropically removed,
	In 2 ml water		remining solution A was added in 100 µl fractions
			untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
			benzene and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
	·		azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours filtered to
	,		yield pale yellow-brown colored solid powder.
213.	Rh(COD)PF6/ S,S chiraphos	Saturated strontium	1 gm davisil was suspended in benzene 25 ml in
	tetrasulfonate 25 mg	chloride solution 2	apparatus described in fig agitated. The temperature of
]	S,S chiraphos tetrasulfonate	ml	the suspension was slowly raised such that it gently
	25 mg		refluxes to which was added 100 µl portion of solution
	Sodium alginate 100 mg		A and solvent component was azeotropically removed,
	dissolved in 2 ml water		remining solution A was added in 100 µl fractions
			untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
		,	benzene and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
·			azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours filtered to
			yield pale yellow colored solid powder.
214.	Rh(COD)PF ₆ / S,S chiraphos	Saturated barium nitrate	1 gm davisil was suspended in benzene 25 ml in apparatus
214.	tetrasulfonate 25 mg	solution 2 ml	described in fig agitated. The temperature of the suspension
	S,S chiraphos tetrasulfonate 25		was slowly raised such that it gently refluxes to which was
	mg		added 100 µl portion of solution A and solvent component was
	Sodium alginate 100 mg dissolved	,	azeotropically removed, remining solution A was added in 100
	in 2 ml water		μl fractions untill uniform suspension volume of suspension
,			was maintained while maintaining volume by pumping benzene
			and solution B was added in equal fractions over a period of 2
	·		hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was
			agitated for 10 hours filtered to yield pale yellow colored solid
		•	powder.
215.	Rh(COD)PF ₆ / S,S chiraphos	Saturated barium	1 gm alumina was suspended in benzene 25 ml in
	tetrasulfonate 25 mg	nitrate solution 2 ml	apparatus described in fig agitated. The temperature of
	S,S chiraphos tetrasulfonate		the suspension was slowly raised such that it gently
]	25 mg		refluxes to which was added 100 µl portion of solution
	Sodium alginate 100 mg		A and solvent component was azeotropically removed,
	dissolved in 2 ml water		remining solution A was added in 100 µl fractions
			untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
			benzene and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
			azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours filtered to
	·		yield pale yellow colored solid powder.

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216.	Rh(COD)PF ₆ / S,S chiraphos	Saturated barium	1 gm titania was suspended in benzene 25 ml in
	tetrasulfonate 25 mg	nitrate solution 2 ml	apparatus described in fig agitated. The temperature of
	S,S chiraphos tetrasulfonate		the suspension was slowly raised such that it gently
	25 mg		refluxes to which was added 100 µl portion of solution
	Sodium alginate 100 mg		A and solvent component was azeotropically removed,
	dissolved in 2 ml water		remining solution A was added in 100 µl fractions
			untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
			benzene and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
	·		azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours filtered to
			yield pale yellow colored solid powder.
217.	HRhCO(TPATS) ₃	500 mg Calcium	1 gm titania was suspended in benzene 25 ml in
	10 mg	chloride solution in	apparatus described in fig agitated. The temperature of
	100 mg TPATS	water 5 ml	the suspension was slowly raised such that it gently
	carboxy methyl cellulose		refluxes to which was added 100 µl portion of solution
	sodium 100 mg in 1 ml water		A and solvent component was azeotropically removed,
	Ü		remining solution A was added in 100 µl fractions
			untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
	·		benzene and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
	:		azeotropic water and suspension was further continued.
	•		Formed suspension was agitated for 10 hours filtered to
			yield pale yellow-green colored solid powder.
218.	HRhCO(TPATS) ₃	Strontium chloride	1 gm alumina was suspended in benzene 25 ml in apparatus
210.	10 mg	saturated solution in	described in fig agitated. The temperature of the suspension
	100 mg TPATS	water 5 ml	was slowly raised such that it gently refluxes to which was
	100 mg 111110		
	carboxy methyl cellulose sodium		added 100 µl portion of solution A and solvent component was
	_		
	carboxy methyl cellulose sodium		added 100 µl portion of solution A and solvent component was
	carboxy methyl cellulose sodium		added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene
	carboxy methyl cellulose sodium		added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2
	carboxy methyl cellulose sodium		added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and
	carboxy methyl cellulose sodium		added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was
	carboxy methyl cellulose sodium		added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and
219.	carboxy methyl cellulose sodium	Barium nitrate	added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for 10 hours filtered to yield pale yellow-green colored
219.	carboxy methyl cellulose sodium 100 mg in 1 ml water	Barium nitrate saturated solution in	added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for 10 hours filtered to yield pale yellow-green colored solid powder.
219.	carboxy methyl cellulose sodium 100 mg in 1 ml water HRhCO(TPATS)3 10 mg		added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for 10 hours filtered to yield pale yellow-green colored solid powder. 1 gm bentonite was suspended in benzene 25 ml in
219.	carboxy methyl cellulose sodium 100 mg in 1 ml water HRhCO(TPATS)3	saturated solution in	added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for 10 hours filtered to yield pale yellow-green colored solid powder. 1 gm bentonite was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of
219.	carboxy methyl cellulose sodium 100 mg in 1 ml water HRhCO(TPATS) ₃ 10 mg 100 mg TPATS	saturated solution in	added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for 10 hours filtered to yield pale yellow-green colored solid powder. 1 gm bentonite was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently
219.	carboxy methyl cellulose sodium 100 mg in 1 ml water HRhCO(TPATS) ₃ 10 mg 100 mg TPATS carboxy methyl cellulose	saturated solution in	added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for 10 hours filtered to yield pale yellow-green colored solid powder. 1 gm bentonite was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed,
219.	carboxy methyl cellulose sodium 100 mg in 1 ml water HRhCO(TPATS) ₃ 10 mg 100 mg TPATS carboxy methyl cellulose	saturated solution in	added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for 10 hours filtered to yield pale yellow-green colored solid powder. 1 gm bentonite was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions
219.	carboxy methyl cellulose sodium 100 mg in 1 ml water HRhCO(TPATS) ₃ 10 mg 100 mg TPATS carboxy methyl cellulose	saturated solution in	added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for 10 hours filtered to yield pale yellow-green colored solid powder. 1 gm bentonite was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was
219.	carboxy methyl cellulose sodium 100 mg in 1 ml water HRhCO(TPATS) ₃ 10 mg 100 mg TPATS carboxy methyl cellulose	saturated solution in	added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for 10 hours filtered to yield pale yellow-green colored solid powder. 1 gm bentonite was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping
219.	carboxy methyl cellulose sodium 100 mg in 1 ml water HRhCO(TPATS) ₃ 10 mg 100 mg TPATS carboxy methyl cellulose	saturated solution in	added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for 10 hours filtered to yield pale yellow-green colored solid powder. 1 gm bentonite was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions
219.	carboxy methyl cellulose sodium 100 mg in 1 ml water HRhCO(TPATS) ₃ 10 mg 100 mg TPATS carboxy methyl cellulose	saturated solution in	added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for 10 hours filtered to yield pale yellow-green colored solid powder. 1 gm bentonite was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of
219.	carboxy methyl cellulose sodium 100 mg in 1 ml water HRhCO(TPATS) ₃ 10 mg 100 mg TPATS carboxy methyl cellulose	saturated solution in	added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for 10 hours filtered to yield pale yellow-green colored solid powder. 1 gm bentonite was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued.
219.	carboxy methyl cellulose sodium 100 mg in 1 ml water HRhCO(TPATS) ₃ 10 mg 100 mg TPATS carboxy methyl cellulose	saturated solution in	added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued. Formed suspension was agitated for 10 hours filtered to yield pale yellow-green colored solid powder. 1 gm bentonite was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of

	T	T	
220.	HRhCO(TPATS) ₃	Strontium chloride	1 gm titania was suspended in benzene 25 ml in
	10 mg	saturated solution in	apparatus described in fig agitated. The temperature of
	100 mg TPATS	water 5 ml	the suspension was slowly raised such that it gently
•	carboxy methyl cellulose		refluxes to which was added 100 µl portion of solution
	sodium 100 mg in 1 ml water		A and solvent component was azeotropically removed,
			remining solution A was added in 100 µl fractions
	, i		untill uniform suspension volume of suspension was
1			maintained while maintaining volume by pumping
			benzene and solution B was added in equal fractions
•			over a period of 2 hours and simultanious removal of
	i .		azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours filtered to
			yield pale yellow-green colored solid powder.
221.	HRhCO(TPATS) ₃	Strontium chloride	1 gm davisil was suspended in benzene 25 ml in
1	10 mg	saturated solution in	apparatus described in fig agitated. The temperature of
	100 mg TPATS	water 5 ml	the suspension was slowly raised such that it gently
	carboxy methyl cellulose		refluxes to which was added 100 µl portion of solution
	sodium 100 mg in 1 ml water		A and solvent component was azeotropically removed,
	Soulding 100 mg in 1 mi water		
	·	,	remining solution A was added in 100 µl fractions
			untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
•	• •		benzene and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
			azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours filtered to
			yield pale yellow-green colored solid powder.
222.	HRhCO(BISBIS) 50 mg	Saturated barium nitrate	2 gm davisil was suspended in benzene 25 ml in apparatus
•	BISBIS 200 mg 200 mg sodium sulfate	solution is 5 ml water	described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was
	dissolved in 2 ml water		added 100 µl portion of solution A and solvent component was
			azeotropically removed, remining solution A was added in 100
		•	µl fractions untill uniform suspension volume of suspension
			was maintained while maintaining volume by pumping benzene
			and solution B was added in equal fractions over a period of 2
			hours and simultanious removal of azeotropic water and
			suspension was further continued. Formed suspension was
			agitated for 10 hours filtered to yield pale yellow colored solid powder.
223.	HRhCO(BISBIS) 50 mg	1 g calcium chloride	2 gm davisil was suspended in benzene 25 ml in
	BISBIS 200 mg	solution In 5 ml	apparatus described in fig agitated. The temperature of
	200 mg polyvinyl sulfonic	water	the suspension was slowly raised such that it gently
	acid dissolved in 2 ml water		refluxes to which was added 100 µl portion of solution
			A and solvent component was azeotropically removed,
			remining solution A was added in 100 µl fractions
-			untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
			l benzene and solution R was added in equal fractions
			benzene and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
			over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued.
			over a period of 2 hours and simultanious removal of

224	HBPCO(BIGBIC) 20 mg	Saturated barium	2 gm titania was suspended in benzene 25 ml in
224.	HRhCO(BISBIS) 50 mg	nitrate solution is 5	apparatus described in fig agitated. The temperature of
	BISBIS 200 mg		
	200 mg polyacrylic acid	ml water	the suspension was slowly raised such that it gently
	sodium salt dissolved in 2 ml		refluxes to which was added 100 µl portion of solution
	water		A and solvent component was azeotropically removed,
			remining solution A was added in 100 μl fractions
			untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
			benzene and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
			azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours filtered to
			yield pale yellow colored solid powder.
225.	HRhCO (BISBIS) 50 mg	Saturated strontium	2 gm alumina was suspended in benzene 25 ml in
	BISBIS 200 mg	chloride solution is 5	apparatus described in fig agitated. The temperature of
	200 mg polyvinyl sulfonic	ml water	the suspension was slowly raised such that it gently
	acid dissolved in 2 ml water		refluxes to which was added 100 µl portion of solution
			A and solvent component was azeotropically removed,
	•		remining solution A was added in 100 µl fractions
	,		untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
			benzene and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
		٠	azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours filtered to
			yield pale yellow colored solid powder.
226.	HRhCO(BISBIS) 50 mg	Saturated barium nitrate	2 gm davisil was suspended in benzene 25 ml in apparatus
	BISBIS 200 mg	solution is 5 ml water	described in fig agitated. The temperature of the suspension
	200 mg polyvinyl sulfonic acid		was slowly raised such that it gently refluxes to which was
	dissolved in 2 ml water		added 100 µl portion of solution A and solvent component was
	•	•	azeotropically removed, remining solution A was added in 100
		٠	µl fractions untill uniform suspension volume of suspension
			was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2
			hours and simultanious removal of azeotropic water and
			suspension was further continued. Formed suspension was
			agitated for 10 hours filtered to yield pale yellow colored solid
,			powder.
227.	HRhCO(BISBIS) 50 mg	Saturated barium	2 gm davisil was suspended in benzene 25 ml in
	BISBIS 200 mg	nitrate solution is 5	apparatus described in fig agitated. The temperature of
	200 mg polyvinyl sulfonic	ml water	the suspension was slowly raised such that it gently
	acid dissolved in 2 ml water		refluxes to which was added 100 µl portion of solution
			A and solvent component was azeotropically removed,
		•	remining solution A was added in 100 µl fractions
			untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
			benzene and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
	,		azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours filtered to
			I more more to to more minima to
			yield pale yellow colored solid powder.

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228.	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm davisil was suspended in benzene 25 ml in
	TPPTS 100 mg	barium nitrate 5 ml	apparatus described in fig agitated. The temperature of
	100 mg sodium alginate		the suspension was slowly raised such that it gently
	Dissolved in 2 ml water and		refluxes to which was added 100 µl portion of solution
	0.5 ml butane diol		A and solvent component was azeotropically removed,
			remining solution A was added in 100 µl fractions
	}		untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
			benzene and solution B was added in equal fractions
	-		over a period of 2 hours and simultanious removal of
			azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours filtered to
	·		yield pale yellow colored solid powder.
229.	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm alumina 2 gm was suspended in benzene 25 ml
	TPPTS 100 mg	barium nitrate 5 ml	in apparatus described in fig agitated. The temperature
1	100 mg oxalic acid sodium		of the suspension was slowly raised such that it gently
	salt.		refluxes to which was added 100 µl portion of solution
	Dissolved in 2 ml water and		A and solvent component was azeotropically removed,
	0.5 ml butane diol		remining solution A was added in 100 µl fractions
			untill uniform suspension volume of suspension was
•			maintained while maintaining volume by pumping
			benzene and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
i			azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours filtered to
			yield pale yellow colored solid powder.
230.	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm davisil was suspended in benzene 25 ml in apparatus
	TPPTS 100 mg	strontium chloride 5 ml	described in fig agitated. The temperature of the suspension
	100 mg citric acid		was slowly raised such that it gently refluxes to which was
	Dissolved in 2 ml water and 0.5		added 100 µl portion of solution A and solvent component was
	ml ethylene glycol		azeotropically removed, remining solution A was added in 100
		•	μl fractions untill uniform suspension volume of suspension
			was maintained while maintaining volume by pumping benzene
			and solution B was added in equal fractions over a period of 2
	·		hours and simultanious removal of azeotropic water and
	.		suspension was further continued. Formed suspension was
			agitated for 10 hours filtered to yield pale yellow colored solid powder.
231.	PtCl ₂ (TPPTS) ₂ 50 mg	Command colution of	2 gm davisil was suspended in benzene 25 ml in
231.	-	Saturated solution of	·
	TPPTS 100 mg	barium nitrate 5 ml	apparatus described in fig agitated. The temperature of
	100 mg polyacrylic acid		the suspension was slowly raised such that it gently
	sodium salt.		refluxes to which was added 100 µl portion of solution
	Dissolved in 2 ml water and		A and solvent component was azeotropically removed,
	0.5 ml butane diol		remining solution A was added in 100 µl fractions
			untill uniform suspension volume of suspension was
·			maintained while maintaining volume by pumping
			benzene and solution B was added in equal fractions
		,	over a period of 2 hours and simultanious removal of
			azeotropic water and suspension was further continued.
			azeotropic water and suspension was further continued. Formed suspension was agitated for 10 hours filtered to
			•

232.	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm shreded asbestos roap 2 gm zirconia was
	TPPTS 300 mg	barium nitrate 5 ml	suspended in benzene 25 ml in apparatus described in
	Dissolved in 2 ml water		fig agitated. The temperature of the suspension was
:			slowly raised such that it gently refluxes to which was
			added 100 µl portion of solution A and solvent
			component was azeotropically removed, remining
			solution A was added in 100 µl fractions untill uniform
		•	suspension volume of suspension was maintained while
			maintaining volume by pumping benzene and solution
			B was added in equal fractions over a period of 2 hours
			and simultanious removal of azeotropic water and
			suspension was further continued. Formed suspension
		•	was agitated for 10 hours filtered to yield pale yellow-
		·	gray colored solid powder.
233.	Cobalt N, N'ethylene bis	Saturated barium nitrate	2 gm davisil was suspended in benzene 25 ml in apparatus
	(salicyldiamine) 5-sulfonato	solution in water 5ml	described in fig agitated. The temperature of the suspension
	sodium 100 mg.		was slowly raised such that it gently refluxes to which was
	Sodium phosphate. 500 mg.		added 100 µl portion of solution A and solvent component was
	In 5 ml water		azeotropically removed, remining solution A was added in 100
			ul fractions until uniform suspension volume of suspension was maintained while maintaining volume by pumping
•			benzene and solution B was added in equal fractions over a
• *		İ	period of 2 hours and simultanious removal of azeotropic water
			and suspension was further continued. Formed suspension was
			agitated for 10 hours filtered to yield pale brown colored solid
	·		powder.
234.	Cobalt N, N'ethylene bis	Saturated barium	2 gm alumina was suspended in benzene 25 ml in
	(salicyldiamine) 5-sulfonato	nitrate solution in	apparatus described in fig agitated. The temperature of
	sodium 100 mg.	water 5ml	the suspension was slowly raised such that it gently
	Sodium silicate 500 mg.		refluxes to which was added 100 µl portion of solution
	In 5 ml water		A and solvent component was azeotropically removed,
			remining solution A was added in 100 μl fractions
			untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
	٠.		benzene.and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
	•		azeotropic water and suspension was further continued.
		·	Formed suspension was agitated for 10 hours filtered to
			yield pale brown colored solid powder.
235.	Cobalt N, N'ethylene bis	Saturated barium	2 gm titania was suspended in benzene 25 ml in
	(salicyldiamine) 5-sulfonato	nitrate solution in	apparatus described in fig agitated. The temperature of
, .	sodium 100 mg.	water 5ml	the suspension was slowly raised such that it gently
	Polyvinyl sulfonate sodium.		refluxes to which was added 100 µl portion of solution
	500 mg.		A and solvent component was azeotropically removed,
•	In 5 ml water		remining solution A was added in 100 µl fractions
	, ·		untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
į			benzene.and solution B was added in equal fractions
l			over a period of 2 hours and simultanious removal of
		.	azeotropic water and suspension was further continued.
·			Formed suspension was agitated for 10 hours filtered to
		ļ	yield pale brown colored solid powder.
			

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236.	Cobalt N, N'ethylene bis	Saturated barium nitrate	2 gm zirconia was suspended in benzene 25 ml in apparatus
	(salicyldiamine) 5-sulfonato	solution in water 5ml	described in fig agitated. The temperature of the suspension
	sodium 100 mg.		was slowly raised such that it gently refluxes to which was
	Polyvinyl sulfonate sodium. 500		added 100 µl portion of solution A and solvent component was
	mg.		azeotropically removed, remining solution A was added in 100
	In 5 ml water		μl fractions untill uniform suspension volume of suspension
			was maintained while maintaining volume by pumping
			benzene.and solution B was added in equal fractions over a
			period of 2 hours and simultanious removal of azeotropic water
		·	and suspension was further continued. Formed suspension was
			agitated for 10 hours filtered to yield pale brown colored solid
			powder.
237.	Cobalt N, N'ethylene bis	2g calcium chloride	2 gm shreded asbestos roap was suspended in benzene 25 ml in
	(salicyldiamine) 5-sulfonato	solution in water 5ml	apparatus described in fig agitated. The temperature of the
	sodium 100 mg.		suspension was slowly raised such that it gently refluxes to
•	Polyvinyl sulfonate sodium. 500		which was added 100 µl portion of solution A and solvent
	mg.		component was azeotropically removed, remining solution A
	In 5 ml water		was added in 100 µl fractions untill uniform suspension volume
	1		,
	· ·		of suspension was maintained while maintaining volume by
			pumping benzene and solution B was added in equal fractions
	İ		over a period of 2 hours and simultanious removal of
			azeotropic water and suspension was further continued. Formed
	·		suspension was agitated for. 10 hours filtered to yield gray
			colored solid powder.
238.	Cobalt (II), 4, 4', 4'',4'''-	Saturated strontium	2 gm shreded asbestos roap was suspended in benzene
236.			
*	tetrasulfopthalocynine 500	chloride in 5 ml	25 ml in apparatus described in fig agitated. The
	mg	water	temperature of the suspension was slowly raised such
	And 500 mg sodium sodium		that it gently refluxes to which was added 100 μ l
	poly vinyl sulfonate in 5 ml		portion of solution A and solvent component was
	water		azeotropically removed, remining solution A was
			added in 100 µl fractions untill uniform suspension
			volume of suspension was maintained while
			·
		•	maintaining volume by pumping benzene.and solution
			B was added in equal fractions over a period of 2 hours
			and simultanious removal of azeotropic water and
			suspension was further continued. Formed suspension
			was agitated for. 10 hours filtered to yield blue-gray
			colored solid powder.
239.	Cobalt (II), 4, 4', 4",4"'-	Saturated barium	2 gm keisulghur was suspended in benzene 25 ml in
	tetrasulfopthalocynine. 500	nitrate in 5 ml water	apparatus described in fig agitated. The temperature of
	mg		the suspension was slowly raised such that it gently
			refluxes to which was added 100 µl portion of solution
	phosphate in 5 ml water		A and solvent component was azeotropically removed,
			A and solvent component was azeotropically removed,
			A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was
			A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping
			A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene.and solution B was added in equal fractions
			A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping
			A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene.and solution B was added in equal fractions
			A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued.
			A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene.and solution B was added in equal fractions over a period of 2 hours and simultanious removal of

			0
240.	Cobalt (II), 4, 4', 4",4"'-	Saturated strontium	2 gm keisulghur was suspended in benzene 25 ml in
	tetrasulfopthalocynine. 500	chloride in 5 ml	apparatus described in fig agitated. The temperature of
	mg	water	the suspension was slowly raised such that it gently
	And 500 mg sodium		refluxes to which was added 100 µl portion of solution
·	phosphate in 5 ml water		A and solvent component was azeotropically removed,
		٠	remining solution A was added in 100 μl fractions
	_		untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
			benzene.and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
		٠	azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours and
-			filtered to yield pale blue colored solid powder
241.	Cobalt (II), 4, 4', 4'',4'''-	500mg. CaCl ₂ in 5 ml	2 gm keisulghur was suspended in benzene 25 ml in
	tetrasulfopthalocynine. 500	water	apparatus described in fig agitated. The temperature of
	mg		the suspension was slowly raised such that it gently
	And 500 mg sodium		refluxes to which was added 100 µl portion of solution
	phosphate in 5 ml water		A and solvent component was azeotropically removed,
			remining solution A was added in 100 µl fractions
			untill uniform suspension volume of suspension was
	-		
			maintained while maintaining volume by pumping
• •			benzene and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
			azeotropic water and suspension was further continued.
		,	Formed suspension was agitated for 10 hours and
	•		filtered to yield pale blue colored solid powder
242.	Copper (II), 4, 4', 4'',4'''-	500mg. CaCl ₂ in 5 ml	2 gm keisulghur was suspended in benzene 25 ml in apparatus
	tetrasulfopthalocynine. 500 mg	water	described in fig agitated. The temperature of the suspension
	And 500 mg sodium sulfate in 5		was slowly raised such that it gently refluxes to which was
	ml water		added 100 µl portion of solution A and solvent component was
		•	azeotropically removed, remining solution A was added in 100
			µl fractions untill uniform suspension volume of suspension
		·	was maintained while maintaining volume by pumping benzene.and solution B was added in equal fractions over a
			period of 2 hours and simultanious removal of azeotropic water
			and suspension was further continued. Formed suspension was
	ĺ		agitated for 10 hours and filtered to yield pale blue colored
			solid powder
243.	Copper (II), 4, 4', 4'',4'''-	Saturated strontium	2 gm keisulghur was suspended in benzene 25 ml in
	tetrasulfopthalocynine. 500	chloride in 5 ml	apparatus described in fig agitated. The temperature of
		water	the suspension was slowly raised such that it gently
	mg	water	
	And 500 mg sodium silicate		refluxes to which was added 100 µl portion of solution
	in 5 ml water		A and solvent component was azeotropically removed,
			remining solution A was added in 100 μl fractions
			remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was
			untill uniform suspension volume of suspension was
			untill uniform suspension volume of suspension was maintained while maintaining volume by pumping
			untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions
			untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene.and solution B was added in equal fractions over a period of 2 hours and simultanious removal of
			untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene.and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued.

	T. C (15) A. A. A. A. A. A. A.	L Commented Landson	2 gm keisulghur was suspended in benzene 25 ml in
244.	Copper (II), 4, 4', 4'',4'''-	Saturated barium	
	tetrasulfopthalocynine. 500	nitrate in 5 ml water	apparatus described in fig agitated. The temperature of
	mg		the suspension was slowly raised such that it gently
1	And 500 mg sodium silicate		refluxes to which was added 100 µl portion of solution
ļ	in 5 ml water	,	A and solvent component was azeotropically removed,
			remining solution A was added in 100 µl fractions
			untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
			benzene.and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
			azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours and
		: .	filtered to yield pale blue colored solid powder.
245.	Copper (II), 4, 4', 4'',4'''-	Saturated barium	2 gm bentonite was suspended in benzene 25 ml in
	tetrasulfopthalocynine. 500	nitrate in 5 ml water	apparatus described in fig agitated. The temperature of
	mg		the suspension was slowly raised such that it gently
	And 500 mg sodium silicate		refluxes to which was added 100 µl portion of solution
	in 5 ml water		A and solvent component was azeotropically removed,
			remining solution A was added in 100 µl fractions
			untill uniform suspension volume of suspension was
	,		maintained while maintaining volume by pumping
			benzene and solution B was added in equal fractions
·			over a period of 2 hours and simultanious removal of
·			azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours and
			filtered to yield pale blue colored solid powder.
246.	Copper (II), 4, 4', 4'',4'''-	Saturated strontiun	2 gm bentonite was suspended in benzene 25 ml in apparatus
2.0.	tetrasulfopthalocynine. 500 mg	chloride in 5 ml water	described in fig agitated. The temperature of the suspension
	And 500 mg sodium silicate in 5		was slowly raised such that it gently refluxes to which was
·	ml water		added 100 µl portion of solution A and solvent component was
			azeotropically removed, remining solution A was added in 100
			μl fractions untill uniform suspension volume of suspension
			was maintained while maintaining volume by pumping
	·		benzene.and solution B was added in equal fractions over a
			period of 2 hours and simultanious removal of azeotropic water
			and suspension was further continued. Formed suspension was
247.	Manganese(II), 4, 4', 4'',4'''-	Saturated strontiun	and suspension was further continued. Formed suspension was agitated for 10 hours and filtered to yield pale blue colored
247.	Manganese(II), 4, 4', 4'',4'''- tetrasulfopthalocynine. 500	Saturated strontiun	and suspension was further continued. Formed suspension was agitated for 10 hours and filtered to yield pale blue colored solid powder.
247.	l - ' 1		and suspension was further continued. Formed suspension was agitated for 10 hours and filtered to yield pale blue colored solid powder. 2 gm davisil was suspended in benzene 25 ml in
247.	tetrasulfopthalocynine. 500	chloride in 5 ml	and suspension was further continued. Formed suspension was agitated for 10 hours and filtered to yield pale blue colored solid powder. 2 gm davisil was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of
247.	tetrasulfopthalocynine. 500	chloride in 5 ml	and suspension was further continued. Formed suspension was agitated for 10 hours and filtered to yield pale blue colored solid powder. 2 gm davisil was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution
247.	tetrasulfopthalocynine. 500 mg And 500 mg sodium silicate	chloride in 5 ml	and suspension was further continued. Formed suspension was agitated for 10 hours and filtered to yield pale blue colored solid powder. 2 gm davisil was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed,
247.	tetrasulfopthalocynine. 500 mg And 500 mg sodium silicate	chloride in 5 ml	and suspension was further continued. Formed suspension was agitated for 10 hours and filtered to yield pale blue colored solid powder. 2 gm davisil was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions
247.	tetrasulfopthalocynine. 500 mg And 500 mg sodium silicate	chloride in 5 ml	and suspension was further continued. Formed suspension was agitated for 10 hours and filtered to yield pale blue colored solid powder. 2 gm davisil was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was
247.	tetrasulfopthalocynine. 500 mg And 500 mg sodium silicate	chloride in 5 ml	and suspension was further continued. Formed suspension was agitated for 10 hours and filtered to yield pale blue colored solid powder. 2 gm davisil was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping
247.	tetrasulfopthalocynine. 500 mg And 500 mg sodium silicate	chloride in 5 ml	and suspension was further continued. Formed suspension was agitated for 10 hours and filtered to yield pale blue colored solid powder. 2 gm davisil was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene.and solution B was added in equal fractions
247.	tetrasulfopthalocynine. 500 mg And 500 mg sodium silicate	chloride in 5 ml	and suspension was further continued. Formed suspension was agitated for 10 hours and filtered to yield pale blue colored solid powder. 2 gm davisil was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of
247.	tetrasulfopthalocynine. 500 mg And 500 mg sodium silicate	chloride in 5 ml	and suspension was further continued. Formed suspension was agitated for 10 hours and filtered to yield pale blue colored solid powder. 2 gm davisil was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of azeotropic water and suspension was further continued.
247.	tetrasulfopthalocynine. 500 mg And 500 mg sodium silicate	chloride in 5 ml	and suspension was further continued. Formed suspension was agitated for 10 hours and filtered to yield pale blue colored solid powder. 2 gm davisil was suspended in benzene 25 ml in apparatus described in fig agitated. The temperature of the suspension was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was azeotropically removed, remining solution A was added in 100 µl fractions untill uniform suspension volume of suspension was maintained while maintaining volume by pumping benzene and solution B was added in equal fractions over a period of 2 hours and simultanious removal of

248.	Manganese(II), 4, 4', 4",4"'-	Saturated barium	2 gm davisil was suspended in benzene 25 ml in
240.	tetrasulfopthalocynine. 500	nitrate in 5 ml water	apparatus described in fig agitated. The temperature of
	mg		the suspension was slowly raised such that it gently
	And 500 mg sodium silicate		refluxes to which was added 100 µl portion of solution
·	in 5 ml water		A and solvent component was azeotropically removed,
	in 5 im water		remining solution A was added in 100 µl fractions
	·		untill uniform suspension volume of suspension was
	·		maintained while maintaining volume by pumping
• •	·		benzene and solution B was added in equal fractions
	,		over a period of 2 hours and simultanious removal of
			azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours and
			filtered to yield pale blue colored solid powder.
249.	Manganese(II), 4, 4', 4",4"'-	Saturated barium	2 gm alumina was suspended in benzene 25 ml in
	tetrasulfopthalocynine. 500	nitrate in 5 ml water	apparatus described in fig agitated. The temperature of
	mg	,	the suspension was slowly raised such that it gently
	And 500 mg sodium silicate		refluxes to which was added 100 µl portion of solution
	in 5 ml water		A and solvent component was azeotropically removed,
			remining solution A was added in 100 µl fractions
-		1	untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
			benzene.and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
			azeotropic water and suspension was further continued.
			Formed suspension was agitated for 10 hours and
·			filtered to yield pale blue colored solid powder.
250.	Manganese(II), 4, 4', 4'',4'''-	Saturated barium nitrate	2 gm alumina was suspended in benzene 25 ml in apparatus
	tetrasulfopthalocynine. 500 mg	in 5 ml water	described in fig agitated. The temperature of the suspension
	And 500 mg sodium polyvinyl		was slowly raised such that it gently refluxes to which was added 100 µl portion of solution A and solvent component was
	sulfonate in 5 ml water	<u> </u>	azeotropically removed, remining solution A was added in 100
			µl fractions untill uniform suspension volume of suspension
•			was maintained while maintaining volume by pumping
		·	benzene.and solution B was added in equal fractions over a
			period of 2 hours and simultanious removal of azeotropic water
,*			and suspension was further continued. Formed suspension was
			agitated for 10 hours and filtered to yield pale blue colored
			solid powder.
251.	Iron (III), 4, 4', 4'',4'''-	Saturated strontium	2 gm davisil was suspended in benzene 25 ml in
·	tetrasulfopthalocynine oxygen	chloride in 5 ml	apparatus described in fig agitated. The temperature of
	adduct. 500 mg	water	the suspension was slowly raised such that it gently
	And 500 mg sodium sulfate in		refluxes to which was added 100 μl portion of solution
	5 ml water		A and solvent component was azeotropically removed,
•		,	remining solution A was added in 100 µl fractions
			untill uniform suspension volume of suspension was
			maintained while maintaining volume by pumping
		1	benzene.and solution B was added in equal fractions
			over a period of 2 hours and simultanious removal of
			azeotropic water and suspension was further continued.
	,		Formed suspension was agitated for 10 hours and
			filtered to yield pale blue colored solid powder.
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252.	Iron (III), 4, 4', 4'',4'''-	Saturated barium	2 gm davisil was suspended in benzene 25
	tetrasulfopthalocynine	nitrate in water 5	ml in apparatus described in fig agitated.
	oxygen adduct. 500 mg	ml	The temperature of the suspension was
	And 500 mg sodium		slowly raised such that it gently refluxes to
	sulfate in 5 ml water		which was added 100 µl portion of solution
			A and solvent component was
			azeotropically removed, remining solution
			A was added in 100 µl fractions untill
			uniform suspension volume of suspension
			was maintained while maintaining volume
			by pumping benzene.and solution B was
			added in equal fractions over a period of 2
			hours and simultanious removal of
			azeotropic water and suspension was further
	•		continued. Formed suspension was agitated
	·		for 10 hours and filtered to yield pale blue
			colored solid powder.
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Examples 253 -336

Preparation of catalytic formulation by deposition precipitation in fluidized bed

The following examples illustrate one of the procedures for the preparation of the catalytic formulation of the invention in accordance with the method of formulation known as co precipitation near the surface of the solid support in fluidized bed.

The general procedure for the preparation of heterogeneous catalytic formulation is described herein as making of a solution of anionically charged catalytic entity, catalytically inert anionic additive (termed as solution A) and solution of group II A metal ions (termed as solution B). The specified amount of support pretreated as described in earlier is charged in fluidization vessel and solids were fluidized with flow of argon. Temperature of the fluidization chamber was raised to specified temperature. Solution A was sprayed over the bed over the specified period of time in such a way that solids donot form lumps. Fluidization was continued further for specified period of time and solution B was similarly sprayed and fluidization was continued for specified period of time. Solids were discharged from vessel and aged for specified time. Catalytic formulation thus

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(4).

formed were washed with water, methanol, diethyl ether and dried in vacuum. Dry powder was stored under argon in gas tight vessel. These solid catalytic formulations can be used for appropriate reactions depending upon catalytically active entity incorporated in it.

Note1: solution A is prepared by dissolving anionic components including anionic complex and additives to make homogeneous solution in degassed solvents. And resulting solution is also degassed by purging argon.

Note 2: solution B is prepared by dissolving group IIA metal salts. Solution was degassed prior to use

Note 3: fluidized bed deposition was carried out in equipment described in figure

Example	Solution A	Solution B	Procedure
253	HRhCO(TPPTS)3, 50 mg, TPPTS 200 mg. 500µl ethylene glycol dissolved in water 2 ml	Saturated barium nitrate in water 2 ml	2 gm Davisil was fluidized in the current of argon and temperature of the fluidization vessel was raised to 50 °C and solution A was sprayed over a period of 2 hours once solids were free flowing solution B was sprayed over 2 hours and fluidization was continued fourther 2 hours. Solids were discharged and aged for 24 hours to yield pale yellow colored solid powder.
254	HRhCO(TPPTS)3, 50 mg, TPPTS 200 mg. 500µl ethylene glycol dissolved in water 2 ml	Saturated strontium chloride in water 2ml	2 gm Davisil was fluidized in the current of argon and temperature of the fluidization vessel was raised to 50 °C and solution A was sprayed over a period of 2 hours once solids were free flowing solution B was sprayed over 2 hours and fluidization was continued fourther 2 hours. Solids were discharged and aged for 24 hours to yield pale yellow colored solid powder.
255	HRhCO(TPPTS)3, 50 mg, TPPTS 200 mg. 500µl ethylene glycol dissolved in water 2 ml	500 mg of calcium chloride in 2 ml water	2 gm Davisil was fluidized in the current of argon and temperature of the fluidization vessel was raised to 50°C and solution A was sprayed over a period of 2 hours once solids were free flowing solution B was sprayed over 2 hours and fluidization was continued fourther 2 hours. Solids were discharged and aged for 24 hours to yield pale yellow colored solid powder.
256	HRhCO(TPPTS)3, 50 mg, TPPTS 200 mg. 500μl ethylene glycol dissolved in water 2 ml	Barium nitrate saturated solution in water	2 gm γ -alumina was fluidized in the current of argon and temperature of the fluidization vessel was raised to 50 °C and solution A was sprayed over a period of 2 hours once solids were free flowing solution B was sprayed over 2 hours and fluidization was continued fourther 2 hours. Solids were discharged and aged for 24 hours to yield pale yellow colored solid powder.

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257.	HRhCO(TPPTS)3, 50 mg,	Strontium chloride	2 gm γ-alumina was fluidized in the current of argon
	TPPTS 200 mg. 500µl	saturated solution in	and temperature of the fluidization vessel was raised to
	ethylene glycol dissolved in	water 2ml	50 °C and solution A was sprayed over a period of 2
	water 2 ml		hours once solids were free flowing solution B was
		•	sprayed over 2 hours and fluidization was continued
·			fourther 2 hours. Solids were discharged and aged for
			24 hours to yield pale yellow colored solid powder.
258.	HRhCO (TPPTS) 50 mg,	Calcium chloride 500	2 gm γ-alumina was was fluidized in the current of
	TPPTS 200 mg. 500μl	mg solution in 2 ml	argon and temperature of the fluidization vessel was
	ethylene glycol dissolved in	water	raised to 50 °C and solution A was sprayed over a
•	water 2 ml		period of 2 hours once solids were free flowing
			solution B was sprayed over 2 hours and fluidization
	,		was continued fourther 2 hours. Solids were discharged
			and aged for 24 hours to yield pale yellow colored solid
			powder.
259.	HRhCO(TPPTS) 50 mg,	Barium nitrate	2 gm was fluidized in the current of argon and
	TPPTS 200 mg. 500µl	saturated solution in	temperature of the fluidization vessel was raised to 50
	ethylene glycol dissolved in	water 2 ml	^o C and solution A was sprayed over a period of 2 hours
	water 2 ml		once solids were free flowing solution B was sprayed
			over 2 hours and fluidization was continued fourther 2
	·		hours. Solids were discharged and aged for 24 hours to
			yield pale yellow colored solid powder.
260.	HRhCO(TPPTS) 50 mg,	Strontium chloride	2 gm bentonite was fluidized in the current of argon
	TPPTS 200 mg. 500µl	saturated solution in	and temperature of the fluidization vessel was raised to
	ethylene glycol dissolved in	water	50 °C and solution A was sprayed over a period of 2
	water 2 ml		hours once solids were free flowing solution B was
			sprayed over 2 hours and fluidization was continued
•			fourther 2 hours. Solids were discharged and aged for
			24 hours to yield pale yellow colored solid powder.
261.	HRhCO(TPPTS) 50 mg,	Calcium chloride 500	2 gm bentonite was fluidized in the current of argon
	TPPTS 200 mg. 500µl	mg solution in 2 ml	and temperature of the fluidization vessel was raised to
	ethylene glycol dissolved in	water	50 °C and solution A was sprayed over a period of 2
	water 2 ml		hours once solids were free flowing solution B was
			sprayed over 2 hours and fluidization was continued
			fourther 2 hours. Solids were discharged and aged for
			24 hours to yield pale yellow colored solid powder.
262.	HRhCO(TPPTS) 50 mg,	Barium nitrate	2 gm charcoal was fluidized in the current of argon and
	TPPTS 200 mg. 500µl	saturated solution in	temperature of the fluidization vessel was raised to 50
	ethylene glycol dissolved in	water	^o C and solution A was sprayed over a period of 2 hours
ĺ	water 2 ml		once solids were free flowing solution B was sprayed
			over 2 hours and fluidization was continued fourther 2
			hours. Solids were discharged and aged for 24 hours to
			yield black colored solid powder.
263.	HRhCO(TPPTS) 50 mg,	Strontium chloride	2 gm charcoal was fluidized in the current of argon and
	TPPTS 200 mg. 500µl	saturated solution in	temperature of the fluidization vessel was raised to 50
	ethylene glycol dissolved in	water	⁰ C and solution A was sprayed over a period of 2 hours
	water 2 ml		once solids were free flowing solution B was sprayed
			over 2 hours and fluidization was continued fourther 2
i			hours. Solids were discharged and aged for 24 hours to
1			yield black colored solid powder.
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264.	HRhCO(TPPTS) 50 mg,	Calcium chloride 500	2 gm charcoal was fluidized in the current of argon and
	TPPTS 200 mg. 500µl	mg solution in 2 ml	temperature of the fluidization vessel was raised to 50
	ethylene glycol dissolved in 2	water	^o C and solution A was sprayed over a period of 2 hours
	ml water	•	once solids were free flowing solution B was sprayed
			over 2 hours and fluidization was continued fourther 2
			hours. Solids were discharged and aged for 24 hours to
			yield black colored solid powder.
265.	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	2 gm Davisil was fluidized in the current of argon and
203.	500μl ethylene glycol	saturated solution in	temperature of the fluidization vessel was raised to 50
	TPPTS 200 mg dissolved in	2 ml water	^o C and solution A was sprayed over a period of 2 hours
	2 ml water		once solids were free flowing solution B was sprayed
	Z IIII Water		over 2 hours and fluidization was continued fourther 2
			hours. Solids were discharged and aged for 24 hours to
	· ·		yield light brown colored solid powder.
266.	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Strontium chloride	2 gm Davisil was wetted with 100 μl portion of solution A
200.	500µl ethylene glycol	saturated solution in 2	and evaporated under vaccume 10 mm Hg with simultanious
	TPPTS 200 mg dissolved in 2 ml	ml water	tumbling remining solution A was added in 100 µl fractions
	water.		and solid was isolated (moisture content ~20 %) this powder
			was added to solution B in equal fractions over a period of 2
			hours and suspension was further agitated for 10 hours filtured
			to yield light brown colored solid powder.
267.	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	2 gm γ-alumina was fluidized in the current of argon
	500µl ethylene glycol	saturated solution in	and temperature of the fluidization vessel was raised to
	TPPTS 200 mg Dissolved in	2 ml water	50 °C and solution A was sprayed over a period of 2
	2 ml water		hours once solids were free flowing solution B was
		•	sprayed over 2 hours and fluidization was continued
	, i		fourther 2 hours. Solids were discharged and aged for
			24 hours to yield light brown colored solid powder.
268.	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Strontium chloride	2 gm γ-alumina was fluidized in the current of argon
	500µl ethylene glycol	saturated solution in	and temperature of the fluidization vessel was raised to
	TPPTS 200 mg dissolved in 2	2 ml water	50 °C and solution A was sprayed over a period of 2
	ml water		hours once solids were free flowing solution B was
			sprayed over 2 hours and fluidization was continued
			fourther 2 hours. Solids were discharged and aged for
	ľ		24 hours to yield light brown colored solid powder.
269.	Ru(H)(Cl)(TPPTS)₃ 50 mg	Strontium chloride	2 gm γ-alumina was fluidized in the current of argon
	TPPTS 200 mg	saturated solution in	and temperature of the fluidization vessel was raised to
	500μl ethylene glycol	2 ml water	50 °C and solution A was sprayed over a period of 2
	Sodium polyvinylsulfonate		hours once solids were free flowing solution B was
	500 mg dissolved in 2 ml		sprayed over 2 hours and fluidization was continued
	water		fourther 2 hours. Solids were discharged and aged for
			24 hours to yield yield light brown colored solid
			24 hours to yield yield light brown colored solid powder.
270.	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	
270.	Ru(H)(Cl)(TPPTS) ₃ 50 mg TPPTS 200 mg	Barium nitrate	powder.
270.			powder. 2 gm γ-alumina was fluidized in the current of argon
270.	TPPTS 200 mg	saturated solution in	powder. 2 gm γ-alumina was fluidized in the current of argon and temperature of the fluidization vessel was raised to
270.	TPPTS 200 mg 500µl ethylene glycol	saturated solution in	powder. 2 gm γ-alumina was fluidized in the current of argon and temperature of the fluidization vessel was raised to 50 °C and solution A was sprayed over a period of 2
270.	TPPTS 200 mg 500µl ethylene glycol Sodium polyvinylsulfonate	saturated solution in	powder. 2 gm γ-alumina was fluidized in the current of argon and temperature of the fluidization vessel was raised to 50 °C and solution A was sprayed over a period of 2 hours once solids were free flowing solution B was

271	Ru(H)(Cl)(TPPTS) ₃ 50 mg TPPTS 200 mg 500µl ethylene glycol Sodium polyvinylsulfonate 500 mg Dissolved in 2 ml water	Barium nitrate saturated solution in 2 ml water	2 gm titania was fluidized in the current of argon and temperature of the fluidization vessel was raised to 50 °C and solution A was sprayed over a period of 2 hours once solids were free flowing solution B was sprayed over 2 hours and fluidization was continued fourther 2 hours solids were discharged and aged for 24 hours to yield light brown colored solid powder
272	Ru(H)(Cl)(TPPTS) ₃ 50 mg 500µl ethylene glycol TPPTS 200 mg Sodium polyvinylsulfonate 500 mg dissolved in 2 ml water	Barium nitrate saturated solution in 2 ml water	2 gm zirconia was fluidized in the current of argon and temperature of the fluidization vessel was raised to 50 °C and solution A was sprayed over a period of 2 hours once solids were free flowing solution B was sprayed over 2 hours and fluidization was continued fourther 2 hours. Solids were discharged and aged for 24 hours to yield light brown colored solid powder
273	Ru(H)(Cl)(TPPTS) ₃ 50 mg TPPTS 200 mg 500µl ethylene glycol Sodium polyvinylsulfonate 500 mg Dissolved in 2 ml water	Barium nitrate saturated solution in 2 ml water	2 gm activated charcoal was fluidized in the current of argon and temperature of the fluidization vessel was raised to 50 °C and solution A was sprayed over a period of 2 hours once solids were free flowing solution B was sprayed over 2 hours and fluidization was continued fourther 2 hours. Solids were discharged and aged for 24 hours to yield black colored solid powder.
274	PdCl ₂ (TPPTS) ₂ 10 mg TPPTS 100 mg 500µl ethylene glycol Poly acrylic acid sodium salt in 5 ml	Barium nitrate saturated solution 5 ml	2 gm shreaded asbestos roap was fluidized in the current of argon and temperature of the fluidization vessel was raised to 50 °C and solution A was sprayed over a period of 2 hours once solids were free flowing solution B was sprayed over 2 hours and fluidization was continued fourther 2 hours. Solids were discharged and aged for 24 hours to yield yellow gray colored solid powder.
275	PdCl ₂ (TPPTS) ₂ 10 mg TPPTS 100 mg 500µl ethylene glycol Poly acrylic acid sodium salt in 5 ml	Strontium chloride saturated solution 5 ml	2 gm shreaded asbestos roap was fluidized in the current of argon and temperature of the fluidization vessel was raised to 50 °C and solution A was sprayed over a period of 2 hours once solids were free flowing solution B was sprayed over 2 hours and fluidization was continued fourther 2 hours. Solids were discharged and aged for 24 hours to yield yellow gray colored solid powder.
276	PdCl ₂ (TPPTS) ₂ 10 mg TPPTS 100 mg 500µl ethylene glycol Poly acrylic acid sodium salt in 5 ml water	500 mg calcium chloride in 5 ml water.	2 gm shreaded asbestos roap was fluidized in the current of argon and temperature of the fluidization vessel was raised to 50 °C and solution A was sprayed over a period of 2 hours once solids were free flowing solution B was sprayed over 2 hours and fluidization was continued fourther 2 hours. Solids were discharged and aged for 24 hours to yield yellow gray colored solid powder.

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d of 2 hours
d of 2 hours was sprayed

284.	PdAc ₂ trio tolyl phosphine	Barium nitrate	2 gm charcoal was fluidized in the current of argon and
•	trisulfonated 25 mg	saturated solution	temperature of the fluidization vessel was raised to 50.
	500µl ethylene glycol	5ml	^o C and solution A was sprayed over a period of 2 hours
	tri ortho tolyl phosphine		once solids were free flowing solution B was sprayed
	trisulfonated 100 mg	•	over 2 hours and fluidization was continued fourther 2
	dissolved in 2 ml water		hours. Solids were discharged and aged for 24 hours to
			yield black colored solid powder.
285.	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated barium	1 gm davisil was fluidized in the current of argon and
	TPPTS 100 mg	nitrate in 2 ml water	temperature of the fluidization vessel was raised to 50
	500µl ethylene glycol		°C and solution A was sprayed over a period of 2 hours
	Sodium carboxy methyl		once solids were free flowing solution B was sprayed
	cellulose 100 mg dissolved in		over 2 hours and fluidization was continued fourther 2
	2 ml		hours. Solids were discharged and aged for 24 hours to
	7 III		yield solid powder of white color with blue tinge.
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286.	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated barium	1 gm alumina was fluidized in the current of argon and
<i>⊾</i> o∪.	TPPTS 100 mg	nitrate in 2 ml water	temperature of the fluidization vessel was raised to 50
	1	maac ii 2 iii waci	^o C and solution A was sprayed over a period of 2 hours
	500µl ethylene glycol		once solids were free flowing solution B was sprayed
	Sodium carboxy methyl		over 2 hours and fluidization was continued fourther 2
	cellulose 100 mg dissolved in		
	2 ml		hours. Solids were discharged and aged for 24 hours to
			yield solid powder of white color with blue tinge.
287.	NiCl₂.(TPPTS)₂ 25 mg	Saturated barium	1 gm zirconia was fluidized in the current of argon and
	500µl ethylene glycol	nitrate in 2 ml water	temperature of the fluidization vessel was raised to 50
	TPPTS 100 mg		°C and solution A was sprayed over a period of 2 hours
	Sodium carboxy methyl		once solids were free flowing solution B was sprayed
	cellulose 100 mg dissolved in		over 2 hours and fluidization was continued fourther 2
	2 ml	*	hours. Solids were discharged and aged for 24 hours to
			yield solid powder of white color with blue tinge
288.			
	NiCl₂.(TPPTS)₂ 25 mg	Saturated strontium	1 gm zirconia was fluidized in the current of argon and
-	500µl ethylene glycol	chloride in 2 ml	temperature of the fluidization vessel was raised to 50
	TPPTS 100 mg	water	⁰ C and solution A was sprayed over a period of 2 hours
	Sodium carboxy methyl		once solids were free flowing solution B was sprayed
	cellulose 100 mg dissolved in		over 2 hours and fluidization was continued fourther 2
	2 mi		hours. Solids were discharged and aged for 24 hours to
	·		yield pale blue colored solid powder.
289.			
	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated strontium	1 gm titania was fluidized in the current of argon and
	500µl ethylene glycol	chloride in 2 ml	temperature of the fluidization vessel was raised to 50
	TPPTS 100 mg	water	⁰ C and solution A was sprayed over a period of 2 hours
	Sodium carboxy methyl		once solids were free flowing solution B was sprayed
	cellulose 100 mg dissolved in		over 2 hours and fluidization was continued fourther 2
	2 ml	•	hours. Solids were discharged and aged for 24 hours to
			yield solid powder of white color with blue tinge
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290.	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated strontium	1 gm asbestos was fluidized in the current of argon and
	500µl ethylene glycol	chloride in 2 ml	temperature of the fluidization vessel was raised to 50
	TPPTS 100 mg	water	^o C and solution A was sprayed over a period of 2 hours
	Sodium carboxy methyl		once solids were free flowing solution B was sprayed
	cellulose 100 mg		over 2 hours and fluidization was continued fourther 2
	Dissolved in 2 ml		hours. Solids were discharged and aged for 24 hours to
			yield pale blue colored solid powder.
291.	(IrClCOD) 5 mg exchanged	Saturated strontium	1 gm davisil was fluidized in the current of argon and
	with TPPTS 100 mg.	chloride in 2 ml	temperature of the fluidization vessel was raised to
	Poly acrylic acid sodium salt	water	50°C and solution A was sprayed over a period of 2
	500μl ethylene glycol		hours once solids were free flowing solution B was
	100 mg dissolved in 2 ml		sprayed over 2 hours and fluidization was continued
	water	. !	fourther 2 hours . Solids were discharged and aged for
·] ,		24 hours to yield pale yellow colored solid powder.
292.	(IrCICOD) 5 mg exchanged	Saturated strontium	1 gm keisulghur was fluidized in the current of argon
	with TPPTS 100 mg.	chloride in 2 ml	and temperature of the fluidization vessel was raised to
	Poly acrylic acid sodium salt	water	50 °C and solution A was sprayed over a period of 2
	500µl ethylene glycol		hours once solids were free flowing solution B was
	100 mg dissolved in 2 ml		sprayed over 2 hours and fluidization was continued
	water		fourther 2 hours. Solids were discharged and aged for
		•	24 hours to yield pale yellow colored solid powder.
293.	(IrClCOD) 5 mg exchanged	Saturated strontium	1 gm bentonite was fluidized in the current of argon
	with TPPTS 100 mg.	chloride in 2 ml	and temperature of the fluidization vessel was raised to
	Poly acrylic acid sodium salt	water	50°C and solution A was sprayed over a period of 2
	500µl ethylene glycol		hours once solids were free flowing solution B was
	100 mg dissolved in 2 ml		sprayed over 2 hours and fluidization was continued
	water		fourther 2 hours. Solids were discharged and aged for
	·		24 hours to yield pale yellow colored solid powder.
294.	(RuCl ₂ COD) 5 mg exchanged	Saturated strontium	1 gm davisil was fluidized in the current of argon and
	with diphenyl phosphino	chloride in 2 ml	temperature of the fluidization vessel was raised to 50
	ethane tetrasulfonate 100 mg.	water	^o C and solution A was sprayed over a period of 2 hours
	Poly acrylic acid sodium salt		once solids were free flowing solution B was sprayed
	500µl ethylene glycol		over 2 hours and fluidization was continued fourther 2
	100 mg dissolvd in 2 ml		hours . Solids were discharged and aged for 24 hours to
	water		yield pale yellow colored solid powder.
295.	(RuCl ₂ COD) 5 mg exchanged	Saturated strontium	1 gm davisil was fluidized in the current of argon and
	with diphenyl phosphino	chloride in 2 ml	temperature of the fluidization vessel was raised to
	ethane tetrasulfonate 100 mg.	water	50°C and solution A was sprayed over a period of 2
	Poly acrylic acid sodium salt		hours once solids were free flowing solution B was
•	100 mg		sprayed over 2 hours and fluidization was continued
	500µl ethylene glycol		fourther 2 hours. Solids were discharged and aged for
	dissolved in 2 ml water		24 hours to yield pale yellow colored solid powder.
296.			
	(RuCl ₂ COD) 5 mg exchanged	500 mg calcium	1 gm davisil was fluidized in the current of argon and
	with diphenyl phosphino	chloride in 2 ml	temperature of the fluidization vessel was raised to 50
	ethane tetrasulfonate 100 mg.	water	°C and solution A was sprayed over a period of 2 hours
	Poly acrylic acid sodium salt		once solids were free flowing solution B was sprayed
	500µl ethylene glycol		over 2 hours and fluidization was continued fourther 2
	100 mg dissolved in 2 ml		hours. Solids were discharged and aged for 24 hours to
	water		yield light brown- yellow colored solid powder.
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297.	Rh(COD)PF6/ S,S chiraphos	Saturated strontium	1 gm davisil was fluidized in the current of argon and
4	tetrasulfonate 25 mg	chloride solution 2	temperature of the fluidization vessel was raised to 50
	S,S chiraphos tetrasulfonate	ml	⁰ C and solution A was sprayed over a period of 2 hours
	25 mg		once solids were free flowing solution B was sprayed
	500μl ethylene glycol		over 2 hours and fluidization was continued fourther 2
•	Sodium alginate 100 mg		hours. Solids were discharged and aged for 24 hours to
	dissolved in 2 ml water		yield pale yellow colored solid powder.
298.	Rh(COD)PF ₆ / S,S chiraphos	Saturated barium	1 gm davisil was fluidized in the current of argon and
250.	tetrasulfonate 25 mg	nitrate solution 2 ml	temperature of the fluidization vessel was raised to 50
	S,S chiraphos tetrasulfonate		^o C and solution A was sprayed over a period of 2 hours
	25 mg		once solids were free flowing solution B was sprayed
	500µl ethylene glycol		over 2 hours and fluidization was continued fourther 2
	Sodium alginate 100 mg		hours. Solids were discharged and aged for 24 hours to
	dissolved in 2 ml water		yield pale yellow colored solid powder.
200	Rh(COD)PF ₆ / S ₁ S chiraphos	Saturated barium	1 gm alumina was fluidized in the current of argon and
299.	tetrasulfonate 25 mg	nitrate solution 2 ml	temperature of the fluidization vessel was raised to 50
	S,S chiraphos tetrasulfonate	mant someth 2 mil	^o C and solution A was sprayed over a period of 2 hours
	25 mg		once solids were free flowing solution B was sprayed
	_		over 2 hours and fluidization was continued fourther 2
	500µl ethylene glycol Sodium alginate 100 mg		hours. Solids were discharged and aged for 24 hours to
	dissolved in 2 ml water		yield pale yellow colored solid powder.
		Saturated barium	1 gm titania was fluidized in the current of argon and
300.	Rh(COD)PF6/ S,S chiraphos	nitrate solution 2 ml	temperature of the fluidization vessel was raised to 50
	tetrasulfonate 25 mg	intrate solution 2 im	^o C and solution A was sprayed over a period of 2 hours
	S,S chiraphos tetrasulfonate		once solids were free flowing solution B was sprayed
	25 mg		over 2 hours and fluidization was continued fourther 2
	500µl ethylene glycol	,	hours. Solids were discharged and aged for 24 hours to
	Sodium alginate 100 mg		yield pale yellow colored solid powder.
	dissolved in 2 ml water	500	1 gm titania was fluidized in the current of argon and
301.	HRhCO(TPATS) ₃	500 mg Calcium	
	10 mg	chloride solution in	temperature of the fluidization vessel was raised to 50
	100 mg TPATS	water 5 ml	^o C and solution A was sprayed over a period of 2 hours
	500µl ethylene glycol		once solids were free flowing solution B was sprayed over 2 hours and fluidization was continued fourther 2
	Carboxy methyl cellulose		hours. Solids were discharged and aged for 24 hours to
	sodium 100 mg dissolved in		,
	1 ml water		yield pale yellow-green colored solid powder.
302.	HRhCO(TPATS) ₃	Strontium chloride	1 gm alumina was fluidized in the current of argon and
	10 mg	saturated solution in	temperature of the fluidization vessel was raised to 50
,	500µl ethylene glycol	water 5 ml	OC and solution A was sprayed over a period of 2 hours
	100 mg TPATS		once solids were free flowing solution B was sprayed
	Carboxy methyl cellulose		over 2 hours and fluidization was continued fourther 2
	sodium 100 mg dissolved in 1		hours. Solids were discharged and aged for 24 hours to
	ml water		yield pale yellow-green colored solid powder.
303.	HRhCO(TPATS)3	Barium nitrate	1 gm bentonite was fluidized in the current of argon
	10 mg	saturated solution in	and temperature of the fluidization vessel was raised to
•	100 mg TPATS	water 5 ml	50 °C and solution A was sprayed over a period of 2
	Carboxy methyl cellulose	·	hours once solids were free flowing solution B was
	500µl ethylene glycol		sprayed over 2 hours and fluidization was continued
	sodium 100 mg dissolved in 1		fourther 2 hours. Solids were discharged and aged for
	ml water		24 hours to yield pale yellow-green colored solid
		·	powder.

	Time company	D	1 gm titania was fluidized in the current of argon and
304.	HRhCO(TPATS) ₃	Strontium chloride	temperature of the fluidization vessel was raised to 50
•	10 mg	saturated solution in	
•	100 mg TPATS	water 5 ml	°C and solution A was sprayed over a period of 2 hours
•	Carboxy methyl cellulose		once solids were free flowing solution B was sprayed
	500µl ethylene glycol		over 2 hours and fluidization was continued fourther 2
	sodium 100 mg dissolved in 1	•	hours. Solids were discharged and aged for 24 hours to
	ml water		yield pale yellow-green colored solid powder.
305.	HRhCO(TPATS)3	Strontium chloride	1 gm davisil was fluidized in the current of argon and
	10 mg	saturated solution in	temperature of the fluidization vessel was raised to 50
•	100 mg TPATS	water 5 ml	^o C and solution A was sprayed over a period of 2 hours
	Carboxy methyl cellulose		once solids were free flowing solution B was sprayed
	500µl ethylene glycol		over 2 hours and fluidization was continued fourther 2
	sodium 100 mg dissolved in 1		hours. Solids were discharged and aged for 24 hours to
	ml water		yield pale yellow-green colored solid powder.
306.	HRhCO(BISBIS) 50 mg	Saturated barium	2 gm davisil was fluidized in the current of argon and
	BISBIS 200 mg	nitrate solution is 5	temperature of the fluidization vessel was raised to 50
	200 mg sodium sulfate	ml water	^o C and solution A was sprayed over a period of 2 hours
-	500µl ethylene glycol	 -	once solids were free flowing solution B was sprayed
	dissolved in 2 ml water		over 2 hours and fluidization was continued fourther 2
	dissolved in 2 iii water		hours. Solids were discharged and aged for 24 hours to
	1.	·	yield pale yellow colored solid powder.
			yield pale yellow colored solid powder.
307.	TITLE CO (DICEPTO) CO		2 devicit was fluidied in the aument of argon and
	HRhCO(BISBIS) 50 mg	1 g calcium chloride	2 gm davisil was fluidized in the current of argon and
	BISBIS 200 mg	solution In 5 ml	temperature of the fluidization vessel was raised to
٠	200 mg polyvinyl sulfonic	water	50°C and solution A was sprayed over a period of 2
	acid		hours once solids were free flowing solution B was
	500µl ethylene glycol		sprayed over 2 hours and fluidization was continued
•	dissolved in 2 ml water		fourther 2 hours. Solids were discharged and aged for
			24 hours to yield pale yellow colored solid powder.
308.			
	HRhCO(BISBIS) 50 mg	Saturated barium	2 gm titania was fluidized in the current of argon and
	BISBIS 200 mg	nitrate solution is 5	temperature of the fluidization vessel was raised to 50
	200 mg polyacrylic acid	mi water	^o C and solution A was sprayed over a period of 2 hours
	sodium salt	,	once solids were free flowing solution B was sprayed
	500µl ethylene glycol		over 2 hours and fluidization was continued fourther 2
	dissolved in 2 ml water		hours. Solids were discharged and aged for 24 hours to
	<u> </u>		yield pale yellow colored solid powder.
•	•		
309.			
	HRhCO(BISBIS) 50 mg	Saturated strontium	2 gm alumina was fluidized in the current of argon and
	BISBIS 200 mg	chloride solution is 5	temperature of the fluidization vessel was raised to 50
	200 mg polyvinyl sulfonic	ml water	°C and solution A was sprayed over a period of 2 hours
	acid		once solids were free flowing solution B was sprayed
	500µl ethylene glycol		over 2 hours and fluidization was continued fourther 2
	' ' ' '		
	dissolved in 7 ml makes		hours. Solids were discharged and agen for 24 hours to
	dissolved in 2 ml water		hours. Solids were discharged and aged for 24 hours to
	dissolved in 2 ml water		yield pale yellow colored solid powder.

			Louis and California in the comment of accom-
310.	HRhCO(BISBIS) 50 mg	Saturated barium	2 gm bentonite was fluidized in the current of argon
·	BISBIS 200 mg	nitrate solution is 5	and temperature of the fluidization vessel was raised to
	200 mg polyvinyl sulfonic	ml water	50 °C and solution A was sprayed over a period of 2
	acid		hours once solids were free flowing solution B was
	500µl ethylene glycol		sprayed over 2 hours and fluidization was continued
	dissolved in 2 ml water		fourther 2 hours. Solids were discharged and aged for
			24 hours to yield pale yellow colored solid powder.
311.	HRhCO(BISBIS) 50 mg	Saturated barium	2 gm davisil was fluidized in the current of argon and
	BISBIS 200 mg	nitrate solution is 5	temperature of the fluidization vessel was raised to 50
	200 mg polyvinyl sulfonic	ml water	°C and solution A was sprayed over a period of 2 hours
	acid .		once solids were free flowing solution B was sprayed
	500µl ethylene glycol		over 2 hours and fluidization was continued fourther 2
,	dissolved in 2 ml water		hours. Solids were discharged and aged for 24 hours to
		4	yield pale yellow-green colored solid powder.
312.	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm davisil was fluidized in the current of argon and
·	TPPTS 100 mg	barium nitrate 5 ml	temperature of the fluidization vessel was raised to 50
•	100 mg sodium alginate		^o C and solution A was sprayed over a period of 2 hours
	dissolved in 2 ml water and		once solids were free flowing solution B was sprayed
1 .	0.5 ml butane diol		over 2 hours and fluidization was continued fourther 2
		·	hours. Solids were discharged and aged for 24 hours to
			yield pale yellow-green colored solid powder.
313.	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm alumina was fluidized in the current of argon and
	TPPTS 100 mg	barium nitrate 5 ml	temperature of the fluidization vessel was raised to 50
	100 mg oxalic acid sodium		^o C and solution A was sprayed over a period of 2 hours
	salt dissolved in 2 ml water		once solids were free flowing solution B was sprayed
	and 0.5 ml butane diol		over 2 hours and fluidization was continued fourther 2
		,	hours. Solids were discharged and aged for 24 hours to
			yield pale yellow-green colored solid powder.
314.	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm davisil was fluidized in the current of argon and
	TPPTS 100 mg	strontium chloride 5	temperature of the fluidization vessel was raised to 50
	100 mg citric acid	ml	^o C and solution A was sprayed over a period of 2 hours
	Dissolved in 2 ml water and		once solids were free flowing solution B was sprayed
	0.5 ml ethylene glycol		over 2 hours and fluidization was continued fourther 2
			hours. Solids were discharged and aged for 24 hours to
	,		yield pale yellow-green colored solid powder.
315.	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm davisil was fluidized in the current of argon and
	TPPTS 100 mg	barium nitrate 5 ml	temperature of the fluidization vessel was raised to 50
	100 mg polyacrylic acid		⁰ C and solution A was sprayed over a period of 2 hours
. •	sodium salt.		once solids were free flowing solution B was sprayed
	Dissolved in 2 ml water and		over 2 hours and fluidization was continued fourther 2
·	0.5 ml butane diol		hours. Solids were discharged and aged for 24 hours to
	,		yield pale yellow-green colored solid powder.
316.	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm shreded asbestos roap was fluidized in the current
	TPPTS 300 mg	barium nitrate 5 ml	of argon and temperature of the fluidization vessel was
	500µl ethylene glycol		raised to 50 °C and solution A was sprayed over a
	Dissolved in 2 ml water		period of 2 hours once solids were free flowing
			solution B was sprayed over 2 hours and fluidization
			was continued fourther 2 hours. Solids were discharged
			and aged for 24 hours to yield pale yellow-green
			colored solid powder.

317.	Cobalt N, N'ethylene bis	Saturated barium	2 gm davisil was fluidized in the current of argon and
	(salicyldiamine) 5-sulfonato	nitrate solution in	temperature of the fluidization vessel was raised to 50
	sodium 100 mg.	water 5ml	°C and solution A was sprayed over a period of 2 hours
	Sodium phosphate. 500 mg.	,	once solids were free flowing solution B was sprayed
	500µl ethylene glycol		over 2 hours and fluidization was continued fourther 2
	dissolved in 5 ml water	•	hours. Solids were discharged and aged for 24 hours to
·			yield pale brown colored solid powder.
318.	Cobalt N, N'ethylene bis	Saturated barium	2 gm alumina was fluidized in the current of argon and
	(salicyldiamine) 5-sulfonato	nitrate solution in	temperature of the fluidization vessel was raised to 50
,	sodium 100 mg.	water 5ml	°C and solution A was sprayed over a period of 2 hours.
	500µl ethylene glycol		once solids were free flowing solution B was sprayed
	Sodium silicate 500 mg		over 2 hours and fluidization was continued fourther 2
·	dissolved in 5 ml water		hours . Solids were discharged and aged for 24 hours to
. •	, assorted in 5 in white		yield pale brown colored solid powder.
319.	Cobalt N. N'ethylene bis	Saturated barium	2 gm titania was fluidized in the current of argon and
J17.	(salicyldiamine) 5-sulfonato	nitrate solution in	temperature of the fluidization vessel was raised to 50
		water 5ml	^o C and solution A was sprayed over a period of 2 hours
,	sodium 100 mg.	water Jilli	once solids were free flowing solution B was sprayed
	500µl ethylene glycol		over 2 hours and fluidization was continued fourther 2
	Polyvinyl sulfonate sodium.		hours. Solids were discharged and aged for 24 hours to
	500 mg dissolved in 5 ml		
	water		yield pale brown colored solid powder.
320.	Cobalt N, N'ethylene bis	Saturated barium	2 gm zirconia was fluidized in the current of argon and
	(salicyldiamine) 5-sulfonato	nitrate solution in	temperature of the fluidization vessel was raised to 50
	sodium 100 mg.	water 5ml	^o C and solution A was sprayed over a period of 2 hours
•	500µl ethylene glycol		once solids were free flowing solution B was sprayed
	Polyvinyl sulfonate sodium.		over 2 hours and fluidization was continued fourther 2
,	500 mg dissolved in 5 ml	•	hours. Solids were discharged and aged for 24 hours to
1	water		yield pale brown colored solid powder.
321.	Cobalt N, N'ethylene bis	2g calcium chloride	2 gm shreded asbestos roap was fluidized in the current
	(salicyldiamine) 5-sulfonato	solution in water 5ml	of argon and temperature of the fluidization vessel was
	sodium 100 mg.	:	raised to 50 °C and solution A was sprayed over a
	500µl ethylene glycol	•	period of 2 hours once solids were free flowing
	Polyvinyl sulfonate sodium.		solution B was sprayed over 2 hours and fluidization
	500 mg dissolved in 5 ml		was continued fourther 2 hours. Solids were discharged
	water		and aged for 24 hours to yield gray colored solid
ļ ·		,	powder.
322.	Cobalt (II), 4, 4', 4'',4'''-	Saturated strontium	2 gm shreded asbestos roap was fluidized in the current of
	tetrasulfopthalocynine. 500	chloride in 5 ml water	argon and temperature of the fluidization vessel was raised to
	mg500µl ethylene glycol and 500		50 °C and solution A was sprayed over a period of 2 hours
	mg sodium sodium poly vinyl		once solids were free flowing solution B was sprayed over 2 hours and fluidization was continued fourther 2 hours solids
	sulfonate in 5 ml water		were discharged and aged for 24 hours to yield blue-gray
			colored solid powder.
323.	Cobalt (II), 4, 4', 4'',4'''-	Saturated barium	2 gm kesilghur was fluidized in the current of argon
<i>3-2-</i> .	tetrasulfopthalocynine. 500	nitrate in 5 ml water	and temperature of the fluidization vessel was raised to
	• •		50 °C and solution A was sprayed over a period of 2
	mg500µl ethylene glycol and		hours once solids were free flowing solution B was
·	500 mg sodium phosphate		sprayed over 2 hours and fluidization was continued
	dissolved in 5 ml water		
			fourther 2 hours. Solids were discharged and aged for
			24 hours to yield pale blue colored solid powder.

324.	Cobalt (II), 4, 4', 4",4"'-	Saturated strontium	2 gm kesilghur was fluidized in the current of argon
	tetrasulfopthalocynine 500	chloride in 5 ml	and temperature of the fluidization vessel was raised to
	mg500µl ethylene glycol and	water	50 °C and solution A was sprayed over a period of 2
	500 mg sodium phosphate		hours once solids were free flowing solution B was
	dissolved in 5 ml water		sprayed over 2 hours and fluidization was continued
			fourther 2 hours. Solids were discharged and aged for
			24 hours to yield pale blue colored solid powder.
325.	Cobalt (II), 4, 4', 4",4"'-	500mg. CaCl ₂ in 5 ml	2 gm kesilghur was fluidized in the current of argon
. •	tetrasulfopthalocynine 500	water	and temperature of the fluidization vessel was raised to
	mg		50 °C and solution A was sprayed over a period of 2
	500µl ethylene glycol and		hours once solids were free flowing solution B was
	500 mg sodium phosphate		sprayed over 2 hours and fluidization was continued
	dissolved in 5 ml water		fourther 2 hours. Solids were discharged and aged for
		•	24 hours to yield pale blue colored solid powder.
326.	Copper (II), 4, 4', 4'',4'''-	500mg. CaCl ₂ in 5 ml	2 gm kesilghur was fluidized in the current of argon
1	tetrasulfopthalocynine. 500	water	and temperature of the fluidization vessel was raised to
	mg		50 °C and solution A was sprayed over a period of 2
	500µl ethylene glycol and		hours once solids were free flowing solution B was
ŀ	500 mg sodium sulfate		sprayed over 2 hours and fluidization was continued
	dissolved in 5 ml water		fourther 2 hours. Solids were discharged and aged for
	dissolved in 5 im water		24 hours to yield pale blue colored solid powder.
327.	Copper (II), 4, 4', 4'',4'''-	Saturated strontium	2 gm kesilghur was fluidized in the current of argon
321.	tetrasulfopthalocynine. 500	chloride in 5 ml	and temperature of the fluidization vessel was raised to
	-	water	50 °C and solution A was sprayed over a period of 2
	mg	Watch	hours once solids were free flowing solution B was
	500 ms sodium silicate		sprayed over 2 hours and fluidization was continued
,	500 mg sodium silicate		fourther 2 hours. Solids were discharged and aged for
;	dissolved in 5 ml water		24 hours to yield pale blue colored solid powder.
220	Copper (II), 4, 4', 4'',4'''-	Saturated barium	2 gm kesilghur was fluidized in the current of argon
328.	,	nitrate in 5 ml water	and temperature of the fluidization vessel was raised to
·	· · · · · · · · · · · · · · · · · · ·	nitrate in 5 mi water	50 °C and solution A was sprayed over a period of 2
•	mg		hours once solids were free flowing solution B was
	500µl ethylene glycol		· ·
	And 500 mg sodium silicate		sprayed over 2 hours and fluidization was continued
	in 5 ml water		fourther 2 hours. Solids were discharged and aged for
		0.4	24 hours to yield pale blue colored solid powder.
329.	Copper (II), 4, 4', 4'',4'''-	Saturated barium	2 gm bentonite was fluidized in the current of argon
	tetrasulfopthalocynine 500µl	nitrate in 5 ml water	and temperature of the fluidization vessel was raised to
	ethylene glycol		50 °C and solution A was sprayed over a period of 2
	adduct. 500 mg and 500 mg		hours once solids were free flowing solution B was
	sodium silicate dissolved in 5		sprayed over 2 hours and fluidization was continued
	ml water		fourther 2 hours. Solids were discharged and aged for
			24 hours to yield pale blue colored solid powder.
330.		·	
	Copper (II), 4, 4', 4'',4'''-	Saturated strontiun	2 gm bentonite was fluidized in the current of argon
	tetrasulfopthalocynine. 500	chloride in 5 ml	and temperature of the fluidization vessel was raised to
	mg 500µl ethylene glycol	water	50 °C and solution A was sprayed over a period of 2
	and 500 mg sodium silicate	,	hours once solids were free flowing solution B was
	dissolved in 5 ml water		sprayed over 2 hours and fluidization was continued
,			fourther 2 hours. Solids were discharged and aged for
		. ;	24 hours to yield pale blue colored solid powder.
			

331.	Manganese(II), 4, 4', 4'',4'''-	Saturated strontiun	2 gm davisil was fluidized in the current of argon and
	tetrasulfopthalocynine. 500	chloride in 5 ml	temperature of the fluidization vessel was raised to 50
	mg 500µl ethylene glycol	water	^o C and solution A was sprayed over a period of 2 hours
·	and 500 mg sodium silicate		once solids were free flowing solution B was sprayed
	dissolved in 5 ml water		over 2 hours and fluidization was continued fourther 2
			hours. Solids were discharged and aged for 24 hours to
	·		yield pale blue colored solid powder.
332.	Manganese(II), 4, 4', 4'',4'''-	Saturated barium	2 gm davisil was fluidized in the current of argon and
:	tetrasulfopthalocynine. 500	nitrate in 5 ml water	temperature of the fluidization vessel was raised to 50
	mg 500µl ethylene glycol and		°C and solution A was sprayed over a period of 2 hours
	500 mg sodium silicate		once solids were free flowing solution B was sprayed
	dissolved in 5 ml water	·	over 2 hours and fluidization was continued fourther 2
			hours. Solids were discharged and aged for 24 hours to
			yield pale blue colored solid powder.
333.	Manganese(II), 4, 4', 4",4"'-	Saturated barium	2 gm alumina was fluidized in the current of argon and
	tetrasulfopthalocynine. 500	nitrate in 5 ml water	temperature of the fluidization vessel was raised to 50
	mg 500µl ethylene glycol and		^o C and solution A was sprayed over a period of 2 hours
	500 mg sodium silicate		once solids were free flowing solution B was sprayed
	dissolved in 5 ml water		over 2 hours and fluidization was continued fourther 2
		,	hours. Solids were discharged and aged for 24 hours to
		•	yield pale blue colored solid powder.
334.	Manganese(II), 4, 4', 4",4"'-	Saturated barium	2 gm alumina was fluidized in the current of argon and
	tetrasulfopthalocynine. 500	nitrate in 5 ml water	temperature of the fluidization vessel was raised to 50
	mg		^o C and solution A was sprayed over a period of 2 hours
	500μl ethylene glycol		once solids were free flowing solution B was sprayed
	And 500 mg sodium		over 2 hours and fluidization was continued fourther 2
,	polyvinyl sulfonate in 5 ml	•	hours. Solids were discharged and aged for 24 hours to
	water	·	yield pale blue colored solid powder.
335.	Iron (III), 4, 4', 4'',4'''-	Saturated strontium	2 gm davisil was fluidized in the current of argon and
	tetrasulfopthalocynine oxygen	chloride in 5 ml	temperature of the fluidization vessel was raised to 50
	adduct. 500 mg, 500µl	water	°C and solution A was sprayed over a period of 2 hours
·	ethylene glycol and 500 mg		once solids were free flowing solution B was sprayed
	sodium sulfate dissolved in 5		over 2 hours and fluidization was continued fourther 2
	ml water		hours. Solids were discharged and aged for 24 hours to
	.1		yield pale blue colored solid powder.
336.	Iron (III), 4, 4', 4'',4'''-	Saturated barium	2 gm davisil was fluidized in the current of argon and
	tetrasulfopthalocynine oxygen	nitrate in water 5 ml	temperature of the fluidization vessel was raised to 50
	adduct 500 mg, 500µl		^o C and solution A was sprayed over a period of 2 hours
	ethylene glycol and 500 mg		once solids were free flowing solution B was sprayed
	sodium sulfate dissolved in 5		over 2 hours and fluidization was continued fourther 2
ļ·	ml water		hours. Solids were discharged and aged for 24 hours to
			yield pale blue colored solid powder.

Examples 337-420

Preparation of catalytic formulation by deposition precipitation in coating pan

The following examples illustrate one of the procedures for the preparation of the catalytic formulation of the invention in accordance with the method of formulation known as co precipitation near the surface of the solid support.

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The general procedure for the preparation of heterogeneous catalytic formulation is described herein as making of a solution of anionically charged catalytic entity, catalytically inert anionic additive (termed as solution A) and solution of group II A metal ions (termed as solution B). The specified amount of support pretreated as described in earlier was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to desired temperature under the flow of argon solution a was sprayed on the bed of solids over a specified period followed by spraying solution B resulting solids were tumbled for specified period of time and dried in vacuum. Solids were washed with water, methanol and diethylether and dried. Dry powder was stored under argon in gas tight vessel. These solid catalytic formulations can be used for appropriate reactions depending upon catalytically active entity incorporated in it.

Note1: solution A is prepared by dissolving anionic components including anionic complex and additives to make homogeneous solution in degassed solvents. And resulting solution is also degassed by purging argon.

Note 2: solution B is prepared by dissolving dissolving group IIA metal salts. Solution was degassed prior to use.

Example	Solution A	Solution B	Procedure
337	HRhCO(TPPTS)3, 50 mg, TPPTS 200 mg. 500µl ethylene glycol dissolved in water 2 ml	Saturated barium nitrate in water 2 ml	2 gm Davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum to yield pale yellow colored solid powder.
338	HRhCO(TPPTS)3, 50 mg, TPPTS 200 mg. 500µl ethylene glycol dissolved in water 2 ml	Saturated strontium chloride in water 2ml	2 gm Davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum to yield pale yellow colored solid powder.
339	HRhCO(TPPTS) 50 mg, TPPTS 200 mg. 500µli ethylene glycol dissolved in water 2 ml	500 mg of calcium chloride in 2 ml water	2 gm Davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum to yield pale yellow colored solid powder.

		·	
340.	HRhCO(TPPTS)3, 50 mg,	Barium nitrate	2 gm γ-alumina was charged in a pan, which was
·	TPPTS 200 mg. 500µl	saturated solution in	subsequently set in to rotation. During this procedure
	ethylene glycol dissolved in	water	solids were tumbled. Temperature of the rotating pan
	water 2 ml		was raised to 70 °C, under the flow of argon solution A
			was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			to yield pale yellow colored solid powder.
341.	. HRhCO(TPPTS)3, 50 mg,	Strontium chloride	2 gm γ-alumina was charged in a pan, which was
•	TPPTS 200 mg. 500µl	saturated solution in	subsequently set in to rotation. During this procedure
	ethylene glycol dissolved in	water 2ml	solids were tumbled. Temperature of the rotating pan
	water 2 ml		was raised to 70 °C, under the flow of argon solution A
			was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
·			were further tumbled for 2 hours and dried in vacuum
	·		to yield pale yellow colored solid powder.
342.	HRhCO(TPPTS)3, 50 mg,	Calcium chloride 500	2 gm γ-alumina was charged in a pan, which was
	TPPTS 200 mg. 500µl	mg solution in 2 ml	subsequently set in to rotation. During this procedure
	ethylene glycol dissolved in	water	solids were tumbled. Temperature of the rotating pan
	water 2 ml		was raised to 70 °C, under the flow of argon solution A
			was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			to yield pale yellow colored solid powder.
343.	HRhCO(TPPTS)3, 50 mg,	Barium nitrate	2 gm bentonite was charged in a pan, which was
į	TPPTS 200 mg. 500µl	saturated solution in	subsequently set in to rotation. During this procedure
ļ ļ	ethylene glycol dissolved in	water 2 ml	solids were tumbled. Temperature of the rotating pan
,	water 2 ml		was raised to 70 °C, under the flow of argon solution A
		,	was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			to yield pale yellow colored solid powder.
344.			
	HRhCO(TPPTS)3, 50 mg,	Strontium chloride	2 gm bentonite was charged in a pan, which was
	TPPTS 200 mg. 500µl	saturated solution in	subsequently set in to rotation. During this procedure
	ethylene glycol dissolved in	water	solids were tumbled. Temperature of the rotating pan
	water 2 ml		was raised to 70 °C, under the flow of argon solution A
			was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
		_ ·	to yield pale yellow colored solid powder.
345.	IIDLOO@DDDC\\$ 50	0-1-ik1t3- 500	2 cm hantonita non abarrad in a con indiate con-
	HRhCO(TPPTS)3, 50 mg,	Calcium chloride 500	2 gm bentonite was charged in a pan which was
	TPPTS 200 mg. 500µl	mg solution in 2 ml	subsequently set in to rotation. During this procedure
	ethylene glycol dissolved in	water	solids were tumbled. Temperature of the rotating pan
	water 2 ml		was raised to 70 °C, under the flow of argon solution A
			was sprayed on the bed of solids over a period of 4
			• •
			hours followed by spraying solution B. resulting solids
			• •

346.	HRhCO(TPPTS)3, 50 mg,	Barium nitrate	2 gm charcoal was charged in a pan, which was
	TPPTS 200 mg. 500µl	saturated solution in	subsequently set in to rotation. During this procedure
	ethylene glycol dissolved in	water	solids were tumbled. Temperature of the rotating pan
	water 2 ml		was raised to 70 °C, under the flow of argon solution A
i	,		was sprayed on the bed of solids over a period of 4
	1		hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			to yield black colored solid powder.
347.	HRbCO(TPPTS)3, 50 mg,	Strontium chloride	2 gm charcoal was charged in a pan, which was
	TPPTS 200 mg. 500µl	saturated solution in	subsequently set in to rotation. During this procedure
	ethylene glycol dissolved in	water	solids were tumbled. Temperature of the rotating pan
	water 2 ml		was raised to 70 °C, under the flow of argon solution A
			was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
·			to yield black colored solid powder.
348.	HRhCO(TPPTS)3, 50 mg,	Calcium chloride 500	2 gm charcoal was charged in a pan, which was
	TPPTS 200 mg. 500µl	mg solution in 2 ml	subsequently set in to rotation. During this procedure
	ethylene glycol dissolved in 2	water	solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A
	ml water		was sprayed on the bed of solids over a period of 4
<u>.</u>			hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			to yield black colored solid powder.
240	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	2 gm Davisil was charged in a pan, which was
349.	500µl ethylene glycol	saturated solution in	subsequently set in to rotation. During this procedure
	TPPTS 200 mg dissolved in 2	2 ml water	solids were tumbled. Temperature of the rotating pan
	ml water		was raised to 70 °C, under the flow of argon solution A
	I III Water		was sprayed on the bed of solids over a period of 4
-			hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
		·	to yield light brown colored solid powder.
350.			
	Ru(H)(Cl)(TPPTS)3 50 mg	Strontium chloride	2 gm Davisil was charged in a pan, which was
	500μl ethylene glycol	saturated solution in	subsequently set in to rotation. During this procedure
	TPPTS 200 mg dissolved in 2	2 ml water	solids were tumbled. Temperature of the rotating pan
	ml water.		was raised to 70 °C, under the flow of argon solution A
			was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
•			were further tumbled for 2 hours and dried in vacuum
			to yield light brown colored solid powder.
351.			
	Ru(H)(Cl)(TPPTS) ₃ 50 mg	Barium nitrate	2 gm γ-alumina was charged in a pan, which was
	500µl ethylene glycol	saturated solution in	subsequently set in to rotation. During this procedure
	TPPTS 200 mg Dissolved in	2 ml water	solids were tumbled. Temperature of the rotating pan
	2 ml water		was raised to 70°C, under the flow of argon solution A
			was sprayed on the bed of solids over a period of 4
		ĺ	hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
	·	·	to yield light brown colored solid powder
			to yield light brown colored solid powder

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352. 353.	Ru(H)(Cl)(TPPTS), 50 mg 500µl ethylene glycol TPPTS 200 mg dissolved in 2 ml water Ru(H)(Cl)(TPPTS), 50 mg TPPTS 200 mg 500µl ethylene glycol Sodium polyvinylsulfonate 500 mg dissolved in 2 ml water	Strontium chloride saturated solution in 2 ml water Strontium chloride saturated solution in 2 ml water	2 gm γ-alumina was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum to yield light brown colored solid powder. 2 gm γ-alumina was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum to yield yield light brown colored solid powder.
354.	Ru(H)(Cl)(TPPTS) ₃ 50 mg TPPTS 200 mg 500µl ethylene glycol Sodium polyvinylsulfonate 500 mg dissolved in 2 ml water	Barium nitrate saturated solution in 2 ml water	2 gm γ-alumina was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum to yield light brown colored solid powder.
355.	Ru(H)(Cl)(TPPTS) ₃ 50 mg TPPTS 200 mg 500µl ethylene glycol Sodium polyvinylsulfonate 500 mg dissolved in 2 ml water	Barium nitrate saturated solution in 2 ml water	2 gm titania was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum to yield light brown colored solid powder.
356.	Ru(H)(Cl)(TPPTS), 50 mg 500µl ethylene glycol TPPTS 200 mg Sodium polyvinylsulfonate 500 mg dissolved in 2 ml water	Barium nitrate saturated solution in 2 ml water	2 gm zirconia was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum to yield light brown colored solid powder.
357.	Ru(H)(Cl)(TPPTS) ₃ 50 mg TPPTS 200 mg 500µl ethylene glycol Sodium polyvinylsulfonate 500 mg dissolved in 2 ml water	Barium nitrate saturated solution in 2 ml water	2 gm activated charcoal was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum to yield black colored solid powder.

358.	PdCl ₂ (TPPTS) ₂ 10 mg	Barium nitrate	2 gm shreaded asbestos roap was charged in a pan,
	TPPTS 100 mg	saturated solution 5	which was subsequently set in to rotation. During this
	500µl ethylene glycol	ml	procedure solids were tumbled. Temperature of the
	Poly acrylic acid sodium salt		rotating pan was raised to 70 °C, under the flow of
	dissolved in 5 ml water		argon solution A was sprayed on the bed of solids over
		•	a period of 4 hours followed by spraying solution B.
			resulting solids were further tumbled for 2 hours and
			dried in vacuum to yield yellow gray colored solid
		·	powder.
359.	PdCl ₂ (TPPTS) ₂ 10 mg	Strontium chloride	2 gm shreaded asbestos roap was charged in a pan,
337.	TPPTS 100 mg	saturated solution 5	which was subsequently set in to rotation. During this
	-	ml	procedure solids were tumbled. Temperature of the
	500µl ethylene glycol		rotating pan was raised to 70 °C, under the flow of
	Poly acrylic acid sodium salt	٠	argon solution A was sprayed on the bed of solids over
	dissolved in 5 ml water		a period of 4 hours followed by spraying solution B.
			resulting solids were further tumbled for 2 hours and
. [dried in vacuum to yield yellow gray colored solid
			powder.
360.	PdCl ₂ (TPPTS) ₂ 10 mg	500 mg calcium	2 gm shreaded asbestos roap was charged in a pan,
·	TPPTS 100 mg	chloride in 5 ml	which was subsequently set in to rotation. During this
	500µl ethylene glycol	water.	procedure solids were tumbled. Temperature of the
	Poly acrylic acid sodium salt		rotating pan was raised to 70 °C, under the flow of
	dissolved in 5 ml water	·	argon solution A was sprayed on the bed of solids over
			a period of 4 hours followed by spraying solution B.
·	•	4 · · · · · · · · · · · · · · · · · · ·	resulting solids were further tumbled for 2 hours and
			dried in vacuum to yield yellow gray colored solid
			powder.
361.	PdAc₂BYPYDS 25 mg	Barium nitrate	2 gm davisil was charged in a pan, which was
	BYPYDS 100 mg	saturated solution	subsequently set in to rotation. During this procedure
	500µl ethylene glycol	5ml	solids were tumbled. Temperature of the rotating pan
	dissolved in 2 ml water		was raised to 70 °C, under the flow of argon solution A
	·		was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			were further tumbled for 2 hours and dried in vacuum to yield loght orange colored solid powder.
362.	PdAc ₂ BYPYDS 25 mg	Strontium chloride	
362.	PdAc ₂ BYPYDS 25 mg BYPYDS 100 mg	Strontium chloride saturated solution	to yield loght orange colored solid powder.
362.	BYPYDS 100 mg		to yield loght orange colored solid powder. 2 gm davisil was charged in a pan, which was
362.	BYPYDS 100 mg 500µl ethylene glycol	saturated solution	to yield loght orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure
362.	BYPYDS 100 mg	saturated solution	to yield loght orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A
362.	BYPYDS 100 mg 500µl ethylene glycol	saturated solution	to yield loght orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan
362.	BYPYDS 100 mg 500µl ethylene glycol	saturated solution	to yield loght orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids
362.	BYPYDS 100 mg 500µl ethylene glycol	saturated solution	to yield loght orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum
	BYPYDS 100 mg 500µl ethylene glycol dissolved in 2 ml water	saturated solution 5ml	to yield loght orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum to yield light orange colored solid powder.
362. 363.	BYPYDS 100 mg 500µl ethylene glycol dissolved in 2 ml water PdAc ₂ BYPYDS 25 mg	saturated solution 5ml 500 mg calcium	to yield loght orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum to yield light orange colored solid powder. 2 gm davisil was charged in a pan, which was
	BYPYDS 100 mg 500µl ethylene glycol dissolved in 2 ml water PdAc ₂ BYPYDS 25 mg BYPYDS 100 mg	saturated solution 5ml 500 mg calcium chloride in 5 ml	to yield loght orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum to yield light orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure
	BYPYDS 100 mg 500µl ethylene glycol dissolved in 2 ml water PdAc ₂ BYPYDS 25 mg BYPYDS 100 mg 500µl ethylene glycol	saturated solution 5ml 500 mg calcium	to yield loght orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum to yield light orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan
	BYPYDS 100 mg 500µl ethylene glycol dissolved in 2 ml water PdAc ₂ BYPYDS 25 mg BYPYDS 100 mg	saturated solution 5ml 500 mg calcium chloride in 5 ml	to yield loght orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum to yield light orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A
	BYPYDS 100 mg 500µl ethylene glycol dissolved in 2 ml water PdAc ₂ BYPYDS 25 mg BYPYDS 100 mg 500µl ethylene glycol	saturated solution 5ml 500 mg calcium chloride in 5 ml	to yield loght orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum to yield light orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4
	BYPYDS 100 mg 500µl ethylene glycol dissolved in 2 ml water PdAc ₂ BYPYDS 25 mg BYPYDS 100 mg 500µl ethylene glycol	saturated solution 5ml 500 mg calcium chloride in 5 ml	to yield loght orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum to yield light orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids
	BYPYDS 100 mg 500µl ethylene glycol dissolved in 2 ml water PdAc ₂ BYPYDS 25 mg BYPYDS 100 mg 500µl ethylene glycol	saturated solution 5ml 500 mg calcium chloride in 5 ml	to yield loght orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids were further tumbled for 2 hours and dried in vacuum to yield light orange colored solid powder. 2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A was sprayed on the bed of solids over a period of 4

	Lau promocos		2 gm bentonite was charged in a pan, which was
364.	PdAc ₂ BYPYDS 25 mg	Barium nitrate	<u> </u>
•	BYPYDS 100 mg	saturated solution	subsequently set in to rotation. During this procedure
	500µl ethylene glycol	5ml	solids were tumbled. Temperature of the rotating pan
	dissolved in 2 ml water		was raised to 70 °C, under the flow of argon solution A
		3	was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			to yield loght orange colored solid powder.
365.	PdAc ₂ tri ortho tolyl	Barium nitrate	2 gm bentonite was charged in a pan, which was
	phosphine trisulfonated 25	saturated solution	subsequently set in to rotation. During this procedure
	, mg	5ml	solids were tumbled. Temperature of the rotating pan
	500μl ethylene glycol		was raised to 70 °C, under the flow of argon solution A
	tri ortho tolyl phosphine		was sprayed on the bed of solids over a period of 4
	trisulfonated 100 mg	·	hours followed by spraying solution B. resulting solids
	dissolved in 2 ml water		were further tumbled for 2 hours and dried in vacuum
	·	•	to yield pale brown colored solid powder.
366.	PdAc ₂ tri ortho tolyl	Strontium chloride	2 gm bentonite was charged in a pan, which was
	phosphine trisulfonated 25	saturated solution	subsequently set in to rotation. During this procedure
•	mg	5ml	solids were tumbled. Temperature of the rotating pan
	500μl ethylene glycol	1	was raised to 70 °C, under the flow of argon solution A
	tri ortho tolyl phosphine		was sprayed on the bed of solids over a period of 4
	trisulfonated 100 mg		hours followed by spraying solution B. resulting solids
	dissolved in 2 ml water		were further tumbled for 2 hours and dried in vacuum
	,		to yield pale yellow brown colored solid powder.
367.	PdAc ₂ tri ortho tolyl	Barium nitrate	2 gm alumina was charged in a pan, which was
	phosphine trisulfonated 25	saturated solution	subsequently set in to rotation. During this procedure
	mg	5ml	solids were tumbled. Temperature of the rotating pan
	500µl ethylene glycol	·	was raised to 70 °C, under the flow of argon solution A
	tri ortho tolyl phosphine		was sprayed on the bed of solids over a period of 4
	trisulfonated 100 mg		hours followed by spraying solution B. resulting solids
	Dissolved in 2 ml water		were further tumbled for 2 hours and dried in vacuum
		·	to yield pale yellow brown colored solid powder.
368.		<u> </u>	
	PdAc ₂ tri ortho tolyl	Barium nitrate	2 gm charcoal was charged in a pan, which was
	phosphine trisulfonated 25	saturated solution	subsequently set in to rotation. During this procedure
	mg	5ml	solids were tumbled. Temperature of the rotating pan
	500µl ethylene glycol		was raised to 70 °C, under the flow of argon solution A
	tri ortho tolyl phosphine		was sprayed on the bed of solids over a period of 4
	trisulfonated 100 mg		hours followed by spraying solution B. resulting solids
	dissolved in 2 ml water		were further tumbled for 2 hours and dried in vacuum
	dissolved in 2 ini water]	to yield black colored solid powder.
369.	 		,
JUF.	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated barium	1 gm davisil was charged in a pan, which was
	TPPTS 100 mg	nitrate in 2 ml water	subsequently set in to rotation. During this procedure
	1		solids were tumbled. Temperature of the rotating pan
	500µl ethylene glycol		was raised to 70 °C, under the flow of argon solution A
	Sodium carboxy methyl	1	The raises to 70 C, under the flow of argon solution A
	1	1	tune entraved on the had of colide over a period of A
	cellulose 100 mg		was sprayed on the bed of solids over a period of 4
	1		hours followed by spraying solution B. resulting solids
÷	cellulose 100 mg		

370.	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated barium	1 gm alumina was charged in a pan, which was
	TPPTS 100 mg	nitrate in 2 ml water	subsequently set in to rotation. During this procedure
	500μl ethylene glycol		solids were tumbled. Temperature of the rotating pan
	Sodium carboxy methyl		was raised to 70 °C, under the flow of argon solution A
	cellulose 100 mg		was sprayed on the bed of solids over a period of 4
	dissolved in 2 ml of water		hours followed by spraying solution B. resulting solids
		İ	were further tumbled for 2 hours and dried in vacuum
			to yield solid powder of white color with blue tinge.
371.	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated barium	1 gm zirconia was charged in a pan, which was
•	500µl ethylene glycol	nitrate in 2 ml water	subsequently set in to rotation. During this procedure
	TPPTS 100 mg	:	solids were tumbled. Temperature of the rotating pan
	Sodium carboxy methyl	·	was raised to 70 °C, under the flow of argon solution A
	cellulose 100 mg		was sprayed on the bed of solids over a period of 4
	dissolved in 2 ml of water		hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
	·		to yield solid powder of white color with blue tinge.
372.	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated strontium	1 gm zirconia was charged in a pan, which was
	500µl ethylene glycol	chloride in 2 ml	subsequently set in to rotation. During this procedure
	TPPTS 100 mg	water	solids were tumbled. Temperature of the rotating pan
	Sodium carboxy methyl		was raised to 70 °C, under the flow of argon solution A
	cellulose 100 mg	· .	was sprayed on the bed of solids over a period of 4
	dissolved in 2 ml of water	·	hours followed by spraying solution B. resulting solids
***		, .	were further tumbled for 2 hours and dried in vacuum
	1	·	to yield solid powder of white color with blue tinge.
373.	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated strontium	1 gm titania was charged in a pan, which was
*	500µl ethylene glycol	chloride in 2 ml	subsequently set in to rotation. During this procedure
	TPPTS 100 mg	water	solids were tumbled. Temperature of the rotating pan
	Sodium carboxy methyl		was raised to 70 °C, under the flow of argon solution A
	cellulose 100 mg		was sprayed on the bed of solids over a period of 4
•	dissolved in 2 ml of water		hours followed by spraying solution B. resulting solids
	·		were further tumbled for 2 hours and dried in vacuum
	•		to yield solid powder of white color with blue tinge.
374.			
	NiCl ₂ .(TPPTS) ₂ 25 mg	Saturated strontium	1 gm asbestos was charged in a pan, which was
	500µl ethylene glycol	chloride in 2 ml	subsequently set in to rotation. During this procedure
	TPPTS 100 mg	water	solids were tumbled. Temperature of the rotating pan
	100 mg Sodium carboxy		was raised to 70 °C, under the flow of argon solution A
	methyl cellulose		was sprayed on the bed of solids over a period of 4
	dissolved in 2 ml		hours followed by spraying solution B. resulting solids
	-		were further tumbled for 2 hours and dried in vacuum
	·		to yield gray blue colored solid powder.
375.			<u> </u>
- 3 - 3	(IrClCOD) 5 mg exchanged	Saturated strontium	1 gm davisil was charged in a pan, which was
·	with TPPTS 100 mg.	chloride in 2 ml	subsequently set in to rotation. During this procedure
	100 mg Poly acrylic acid	water	solids were tumbled. Temperature of the rotating pan
	sodium salt 500µl ethylene		was raised to 70 °C, under the flow of argon solution A
	glycol		was sprayed on the bed of solids over a period of 4
	dissolved in 2 ml water		hours followed by spraying solution B. resulting solids
	GISSOIVEG III Z IIII WALEI		, , , ,
			were juriner mindled for 2 notice and direct in vacility is
			were further tumbled for 2 hours and dried in vacuum to yield pale yellow colored solid powder.

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376.	(IrClCOD) 5 mg exchanged	Saturated strontium	1 gm keisulghur was charged in a pan, which was
	with TPPTS 100 mg.	chloride in 2 ml	subsequently set in to rotation. During this procedure
	100 mg Poly acrylic acid	water	solids were tumbled. Temperature of the rotating pan
	sodium salt 500µl ethylene		was raised to 70 °C, under the flow of argon solution A
	glycol		was sprayed on the bed of solids over a period of 4
	dissolved in 2 ml water		hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			to yield pale yellow colored solid powder.
377.	(IrClCOD) 5 mg exchanged	Saturated strontium	1 gm bentonite was charged in a pan, which was
	with TPPTS 100 mg.	chloride in 2 ml	subsequently set in to rotation. During this procedure
	Poly acrylic acid sodium salt	water	solids were tumbled. Temperature of the rotating pan
	500µl ethylene glycol		was raised to 70 °C, under the flow of argon solution A
-	100 mg dissolved in 2 ml		was sprayed on the bed of solids over a period of 4
	water		hours followed by spraying solution B. resulting solids
	· ·		were further tumbled for 2 hours and dried in vacuum
·			to yield pale yellow colored solid powder.
378.	(RuCl ₂ COD) 5 mg exchanged	Saturated strontium	1 gm davisil was charged in a pan, which was
	with diphenyl phosphino	chloride in 2 ml	subsequently set in to rotation. During this procedure
	ethane tetrasulfonate 100 mg.	water	solids were tumbled. Temperature of the rotating pan
	100 mg Poly acrylic acid		was raised to 70 °C, under the flow of argon solution A
	sodium salt 500µl ethylene		was sprayed on the bed of solids over a period of 4
	glycol dissolved in 2 ml water		hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
0.50	(C) (C) (C) (C) (C) (C) (C) (C) (C) (C)	6.4	to yield pale yellow colored solid powder.
379.	(RuCl ₂ COD) 5 mg exchanged	Saturated strontium chloride in 2 ml	1 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure
	with diphenyl phosphino ethane tetrasulfonate 100 mg.	water	solids were tumbled. Temperature of the rotating pan
	100 mg Poly acrylic acid	water .	was raised to 70 °C, under the flow of argon solution A
	sodium salt, 500µl ethylene		was sprayed on the bed of solids over a period of 4
	glycol dissolved in 2 ml		hours followed by spraying solution B. resulting solids
	water		were further tumbled for 2 hours and dried in vacuum
	William	٠.	to yield pale yellow colored solid powder.
380.			
	(RuCl ₂ COD) 5 mg exchanged	500 mg calcium	1 gm davisil was charged in a pan, which was
	with diphenyl phosphino	chloride in 2 ml	subsequently set in to rotation. During this procedure
	ethane tetrasulfonate 100 mg.	water	solids were tumbled. Temperature of the rotating pan
	100 mg Poly acrylic acid		was raised to 70 °C, under the flow of argon solution A
	sodium salt, 500µl ethylene		was sprayed on the bed of solids over a period of 4
	glycol dissolved in 2 ml water		hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			to yield light brown- yellow colored solid powder.
381.			
	Rh(COD)PF S,S chiraphos	Saturated strontium	1 gm davisil was charged in a pan, which was
	tetrasulfonate 25 mg	chloride solution 2	subsequently set in to rotation. During this procedure
	S,S chiraphos	ml	solids were tumbled. Temperature of the rotating pan
	tetrasulfonate 25 mg,		was raised to 70 °C, under the flow of argon solution A
			was sprayed on the bed of solids over a period of 4
	500µl ethylene glycol,	:	hours followed by spraying solution B. resulting solids
:	Sodium alginate 100 mg		were further tumbled for 2 hours and dried in vacuum
	dissolved in 2 ml water		to yield pale yellow colored solid powder.
	<u>. </u>		

382.	Rh(COD)PF ₆ / S,S chiraphos	Saturated barium	1 gm davisil was charged in a pan, which was
	tetrasulfonate 25 mg	nitrate solution 2 ml	subsequently set in to rotation. During this procedure.
	S,S chiraphos tetrasulfonate		solids were tumbled. Temperature of the rotating pan
	25 mg		was raised to 70 °C, under the flow of argon solution A
	500µl ethylene glycol,		was sprayed on the bed of solids over a period of 4
,	Sodium alginate 100 mg,	,*	hours followed by spraying solution B. resulting solids
	dissolved in 2 ml water		were further tumbled for 2 hours and dried in vacuum
		_	to yield pale yellow colored solid powder.
383.	Rh(COD)PF6/ S,S chiraphos	Saturated barium	1 gm alumina was charged in a pan, which was
• • •	tetrasulfonate 25 mg	nitrate solution 2 ml	subsequently set in to rotation. During this procedure
•	S,S chiraphos tetrasulfonate		solids were tumbled. Temperature of the rotating pan
	25 mg, 500µl ethylene glycol,	,	was raised to 70 °C, under the flow of argon solution A
	Sodium alginate 100 mg		was sprayed on the bed of solids over a period of 4
	dissolved in 2 ml water	•	hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			to yield pale yellow colored solid powder.
384.	Rh(COD)PF6/ S,S chiraphos	Saturated barium	1 gm titania was charged in a pan, which was
	tetrasulfonate 25 mg	nitrate solution 2 ml	subsequently set in to rotation. During this procedure
	S,S chiraphos tetrasulfonate		solids were tumbled. Temperature of the rotating pan
	25 mg, 500µl ethylene glycol,		was raised to 70 °C, under the flow of argon solution A
	Sodium alginate 100 mg		was sprayed on the bed of solids over a period of 4
	dissolved in 2 ml water		hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
		·	to yield pale yellow colored solid powder.
385.	HRhCO(TPATS)3	500 mg Calcium	1 gm titania was charged in a pan, which was
	10 mg	chloride solution in	subsequently set in to rotation. During this procedure
٠	100 mg TPATS	water 5 ml	solids were tumbled. Temperature of the rotating pan
	500µl ethylene glycol		was raised to 70 °C, under the flow of argon solution A
	Carboxy methyl cellulose	•	was sprayed on the bed of solids over a period of 4
	sodium 100 mg dissolved in 1		hours followed by spraying solution B. resulting solids
	ml water		were further tumbled for 2 hours and dried in vacuum
			to yield pale yellow-green colored solid powder.
		•	,
386.	HRhCO(TPATS) ₃	Strontium chloride	1 gm alumina was charged in a pan, which was
•	10 mg	saturated solution in	subsequently set in to rotation. During this procedure
	500µl ethylene glycol	water 5 ml	solids were tumbled. Temperature of the rotating pan
	· 100 mg TPATS		was raised to 70 °C, under the flow of argon solution A
	carboxy methyl cellulose		was sprayed on the bed of solids over a period of 4
	sodium 100 mg dissolved in		hours followed by spraying solution B. resulting solids
	1 ml water		were further tumbled for 2 hours and dried in vacuum
		·	to yield pale yellow-green colored solid powder.
387.			
	HRhCO(TPATS)3	Barium nitrate	1 gm bentonite was charged in a pan, which was
	10 mg	saturated solution in	subsequently set in to rotation. During this procedure
	100 mg TPATS	water 5 ml	solids were tumbled. Temperature of the rotating pan
	carboxy methyl cellulose		was raised to 70 °C, under the flow of argon solution A
	500µl ethylene glycol		was sprayed on the bed of solids over a period of 4
	sodium 100 mg dissolved in 1		hours followed by spraying solution B. resulting solids
	ml water		were further tumbled for 2 hours and dried in vacuum
			to yield pale yellow-green colored solid powder.
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388.	HRhCO(TPATS)3	Strontium chloride	1 gm titania was charged in a pan, which was
	10 mg	saturated solution in	subsequently set in to rotation. During this procedure
	100 mg TPATS	water 5 ml	solids were tumbled. Temperature of the rotating pan
	Carboxy methyl cellulose	·	was raised to 70 °C, under the flow of argon solution A
,	500µl ethylene glycol		was sprayed on the bed of solids over a period of 4
	sodium 100 mg dissolved in 1		hours followed by spraying solution B. resulting solids
	ml water		were further tumbled for 2 hours and dried in vacuum
			to yield pale yellow-green colored solid powder.
389.	HRhCO(TPATS)3	Strontium chloride	1 gm davisil was charged in a pan, which was
	10 mg	saturated solution in	subsequently set in to rotation. During this procedure
	100 mg TPATS, sodium	water 5 ml	solids were tumbled. Temperature of the rotating pan
	carboxy methyl cellulose 100		was raised to 70 °C, under the flow of argon solution A
	mg, 500µl ethylene glycol,		was sprayed on the bed of solids over a period of 4
	dissolved in 1 ml water		hours followed by spraying solution B. resulting solids
	· .		were further tumbled for 2 hours and dried in vacuum
			to yield pale yellow-green colored solid powder.
390.	HRhCO(BISBIS) 50 mg	Saturated barium	2 gm davisila was charged in a pan, which was
	BISBIS 200 mg,	nitrate solution is 5	subsequently set in to rotation. During this procedure
	200 mg sodium sulfate,	ml water	solids were tumbled. Temperature of the rotating pan
	500μl ethylene glycol,		was raised to 70 °C, under the flow of argon solution A
	dissolved in 2 ml water		was sprayed on the bed of solids over a period of 4
		: '	hours followed by spraying solution B. resulting solids
		. •	were further tumbled for 2 hours and dried in vacuum
		4 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	to yield pale yellow colored solid powder.
391.	HRhCO(BISBIS) 50 mg	1 g calcium chloride	2 gm davisil was charged in a pan, which was subsequently set in to rotation. During this procedure
·,	BISBIS 200 mg	solution in 5 ml	solids were tumbled. Temperature of the rotating pan
·	200 mg polyvinyl sulfonic	water	was raised to 70 °C, under the flow of argon solution A
	acid, 500µl ethylene glycol,		was sprayed on the bed of solids over a period of 4
	dissolved in 2 ml water	٠	hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
	٠		to yield pale yellow colored solid powder.
392.			
392.	HRhCO(BISBIS) 50 mg	Saturated barium	2 gm titania was charged in a pan, which was
	BISBIS 200 mg	nitrate solution is 5	subsequently set in to rotation. During this procedure
-	200 mg polyacrylic acid	ml water	solids were tumbled. Temperature of the rotating pan
	sodium salt, 500µl ethylene		was raised to 70 °C, under the flow of argon solution A
•	glycol dissolved in 2 ml water		was sprayed on the bed of solids over a period of 4
	gryoor easserver as 2 as well		hours followed by spraying solution B. resulting solids
c			were further tumbled for 2 hours and dried in vacuum
,			to yield pale yellow colored solid powder.
393.	<u> </u>		
- 7	HRhCO(BISBIS) 50 mg	Saturated strontium	2 gm alumina was charged in a pan, which was
	BISBIS 200 mg	chloride solution is 5	subsequently set in to rotation. During this procedure
	200 mg polyvinyl sulfonic	ml water	solids were tumbled. Temperature of the rotating pan
	acid, 500µl ethylene glycol,		was raised to 70 °C, under the flow of argon solution A
	dissolved in 2 ml water		was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			to yield pale yellow colored solid powder.

394.	HRhCO(BISBIS) 50 mg	Saturated barium	2 gm bentonite was charged in a pan, which was
	BISBIS 200 mg, 200 mg	nitrate solution is 5	subsequently set in to rotation. During this procedure
	polyvinyl sulfonic acid, 500µl	ml water	solids were tumbled. Temperature of the rotating pan
•	ethylene glycol, dissolved in		was raised to 70 °C, under the flow of argon solution A
	2 ml water		was sprayed on the bed of solids over a period of 4
,			hours followed by spraying solution B. resulting solids
	·	,	were further tumbled for 2 hours and dried in vacuum
			to yield pale yellow colored solid powder.
395.	HRhCO(BISBIS) 50 mg	Saturated barium	2 gm Davisil was charged in a pan, which was
	BISBIS 200 mg, 200 mg	nitrate solution is 5	subsequently set in to rotation. During this procedure
	polyvinyl sulfonic acid, 500µl	ml water	solids were tumbled. Temperature of the rotating pan
	ethylene glycol, dissolved in		was raised to 70 °C, under the flow of argon solution A
	2 ml water		was sprayed on the bed of solids over a period of 4
·			hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			to yield pale yellow-green colored solid powder.
396.	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm davisil was charged in a pan, which was
	TPPTS 100 mg, 100 mg	barium nitrate 5 ml	subsequently set in to rotation. During this procedure
·	sodium alginate dissolved in		solids were tumbled. Temperature of the rotating pan
	2 ml water and 0.5 ml butane		was raised to 70 °C, under the flow of argon solution A
	diol		was sprayed on the bed of solids over a period of 4
٠	•		hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
•			to yield pale yellow-green colored solid powder.
397.	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm alumina was charged in a pan, which was
	TPPTS 100 mg, 100 mg	barium nitrate 5 ml	subsequently set in to rotation. During this procedure
	oxalic acid sodium salt,		solids were tumbled. Temperature of the rotating pan
	dissolved in 2 ml water and		was raised to 70 °C, under the flow of argon solution A
	0.5 ml butane diol	-	was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
		•	were further tumbled for 2 hours and dried in vacuum
			to yield pale yellow-green colored solid powder.
398.			
	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm davisil was charged in a pan, which was
	TPPTS 100 mg, 100 mg citric	strontium chloride 5	subsequently set in to rotation. During this procedure
	acid, dissolved in 2 ml water	ml	solids were tumbled. Temperature of the rotating pan
	and 0.5 ml ethylene glycol		was raised to 70 °C, under the flow of argon solution A
			was sprayed on the bed of solids over a period of 4
·			hours followed by spraying solution B. resulting solids
	•		were further tumbled for 2 hours and dried in vacuum
			to yield pale yellow-green colored solid powder.
399.			
	PtCl ₂ (TPPTS) ₂ 50 mg	Saturated solution of	2 gm davisil was charged in a pan, which was
	TPPTS 100 mg, 100 mg	barium nitrate 5 ml	subsequently set in to rotation. During this procedure
·	polyacrylic acid sodium salt,		solids were tumbled. Temperature of the rotating pan
	dissolved in 2 ml water and		was raised to 70 °C, under the flow of argon solution A
	0.5 ml butane diol	i	was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
ŀ	l l		to yield pale yellow-green colored solid powder.

400.	PtCl ₂ (TPPTS) ₂ 50 mg, TPPTS	Saturated solution of	2 gm shreded asbestos roap was charged in a pan,
•	300 mg, 500µl ethylene	barium nitrate 5 ml	which was subsequently set in to rotation. During this
	glycol, Dissolved in 2 ml		procedure solids were tumbled. Temperature of the
	water		rotating pan was raised to 70 °C, under the flow of
	}		argon solution A was sprayed on the bed of solids over
	,	·	a period of 4 hours followed by spraying solution B.
			resulting solids were further tumbled for 2 hours and
			dried in vacuum to yield pale yellow-green colored
_			solid powder.
401.	Cobalt N, N'ethylene bis	Saturated barium	2 gm davisil was charged in a pan, which was
	(salicyldiamine) 5-sulfonato	nitrate solution in	subsequently set in to rotation. During this procedure
	sodium 100 mg, Sodium	water 5ml	solids were tumbled. Temperature of the rotating pan
	phosphate. 500 mg.,500µl	, i	was raised to 70 °C, under the flow of argon solution A
	ethylene glycol dissolved in 5	•	was sprayed on the bed of solids over a period of 4
	ml water		hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			to yield pale brown colored solid powder.
402.	Cobalt N, N'ethylene bis	Saturated barium	2 gm alumina was charged in a pan, which was
776.	(salicyldiamine) 5-sulfonato	nitrate solution in	subsequently set in to rotation. During this procedure
,	sodium 100 mg, 500µl	water 5ml	solids were tumbled. Temperature of the rotating pan
	ethylene glycol, Sodium	water Jan	was raised to 70 °C, under the flow of argon solution A
	silicate 500 mg dissolved in 5		was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
	ml water	. •	were further tumbled for 2 hours and dried in vacuum
		•	to yield pale brown colored solid powder.
403.	Cobalt N, N'ethylene bis	Saturated barium	2 gm titania was charged in a pan, which was
403.		nitrate solution in	subsequently set in to rotation. During this procedure
	(salicyldiamine) 5-sulfonato	water 5ml	solids were tumbled. Temperature of the rotating pan
	sodium 100 mg,500µl	water Jun	was raised to 70 °C, under the flow of argon solution A
٠.	ethylene glycol, Polyvinyl		was sprayed on the bed of solids over a period of 4
	sulfonate sodium. 500 mg	• ,	hours followed by spraying solution B. resulting solids
,	dissolved in 5 ml water		were further tumbled for 2 hours and dried in vacuum
			to yield pale brown colored solid powder.
404.	Cobalt N, N'ethylene bis	Saturated barium	2 gm zirconia was charged in a pan, which was
	(salicyldiamine) 5-sulfonato	nitrate solution in	subsequently set in to rotation. During this procedure
	sodium 100 mg, 500µl	water 5ml	solids were tumbled. Temperature of the rotating pan
	ethylene glycol, polyvinyl		was raised to 70 °C, under the flow of argon solution A
	sulfonaté sodium. 500 mg		was sprayed on the bed of solids over a period of 4
•	dissolved in 5 ml water		hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			to yield pale brown colored solid powder.
405.	Cobalt N, N'ethylene bis	2g calcium chloride	2 gm shreded asbestos roap was charged in a pan,
•	(salicyldiamine) 5-sulfonato	solution in water 5ml	which was subsequently set in to rotation. During this
	sodium 100 mg, 500µl	,	procedure solids were tumbled. Temperature of the
	ethylene glycol, polyvinyl	•	rotating pan was raised to 70 °C, under the flow of
	sulfonate sodium. 500 mg in		argon solution A was sprayed on the bed of solids over
	5 ml water	•	a period of 4 hours followed by spraying solution B.
			resulting solids were further tumbled for 2 hours and
			dried in vacuum to yield gray colored solid powder.
	1	لب ــــــــــــــــــــــــــــــــــــ	

406.	Caballe (TD 4 4) 41/41/1	Saturated strontium	2 gm shreded asbestos roap was charged in a pan,
406.	Cobalt (II), 4, 4', 4'',4'''-	chloride in 5 ml	which was subsequently set in to rotation. During this
<i>.</i> ·	tetrasulfopthalocynine 500		procedure solids were tumbled. Temperature of the
	mg500µl ethylene glycol and	water	rotating pan was raised to 70 °C, under the flow of
	500 mg sodium sodium poly		argon solution A was sprayed on the bed of solids over
	vinyl sulfonate in 5 ml water		
·			a period of 4 hours followed by spraying solution B.
			resulting solids were further tumbled for 2 hours and
			dried in vacuum to yield blue-gray colored solid
	,		powder.
407.	Cobalt (II), 4, 4', 4",4"'-	Saturated barium	2 gm kesilghur was charged in a pan, which was
	tetrasulfopthalocynine 500	nitrate in 5 ml water	subsequently set in to rotation. During this procedure
	mg500µl ethylene glycol and		solids were tumbled. Temperature of the rotating pan
	500 mg sodium phosphate in		was raised to 70 °C, under the flow of argon solution A
	5 ml water		was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
•			were further tumbled for 2 hours and dried in vacuum
			to yield pale blue colored solid powder.
408.	Cobalt (II), 4, 4', 4'',4'''-	Saturated strontium	2 gm kesilghur was charged in a pan, which was
	tetrasulfopthalocynine 500	chloride in 5 ml	subsequently set in to rotation. During this procedure
	mg500µl ethylene glycol and	water	solids were tumbled. Temperature of the rotating pan
	500 mg sodium phosphate in	•	was raised to 70 °C, under the flow of argon solution A
•	5 ml water		was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
	•	•	to yield pale blue colored solid powder.
409.	Cobalt (II), 4, 4', 4",4"'-	500mg. CaCl ₂ in 5 ml	2 gm kesilghur was charged in a pan, which was
	tetrasulfopthalocynine 500	water	subsequently set in to rotation. During this procedure
	mg, 500µl ethylene glycol		solids were tumbled. Temperature of the rotating pan
	and 500 mg sodium		was raised to 70 °C, under the flow of argon solution A
	phosphate in 5 ml water		was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			to yield pale blue colored solid powder.
410.		•	
	Copper (II), 4, 4', 4'',4'''-	500mg. CaCl ₂ in 5 ml	2 gm kesilghur was charged in a pan, which was
•	tetrasulfopthalocynine. 500	water	subsequently set in to rotation. During this procedure
	mg, 500µl ethylene glycol		solids were tumbled. Temperature of the rotating pan
	and 500 mg sodium sulfate in		was raised to 70 °C, under the flow of argon solution A
	5 ml water		was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
-		II	were further tumbled for 2 hours and dried in vacuum
			to yield pale blue colored solid powder.
411.	Copper (II), 4, 4', 4'',4'''-	Saturated strontium	2 gm kesilghur was charged in a pan, which was
	tetrasulfopthalocynine. 500	chloride in 5 ml	subsequently set in to rotation. During this procedure
	mg, 500µl ethylene glycol	water	solids were tumbled. Temperature of the rotating pan
·	and 500 mg sodium silicate in		was raised to 70 $^{\circ}$ C, under the flow of argon solution A
	5 ml water		was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			to yield pale blue colored solid powder.
	<u> </u>		<u> </u>

	·		
412.	Copper (II), 4, 4', 4",4"'-	Saturated barium	2 gm kesilghur was charged in a pan, which was
·	tetrasulfopthalocynine. 500	nitrate in 5 ml water	subsequently set in to rotation. During this procedure
	mg, 500µl ethylene glycol		solids were tumbled. Temperature of the rotating pan
. 1	and 500 mg sodium silicate in	· ·	was raised to 70 °C, under the flow of argon solution A
	5 ml water		was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			to yield pale blue colored solid powder.
413.	Copper (II), 4, 4', 4",4"'-	Saturated barium	2 gm bentonite was charged in a pan, which was
	tetrasulfopthalocynine 500µl	nitrate in 5 ml water	subsequently set in to rotation. During this procedure
	ethylene glycol, 500 mg and		solids were tumbled. Temperature of the rotating pan
ŀ	500 mg sodium silicate in 5		was raised to 70 °C, under the flow of argon solution A
	ml water	•	was sprayed on the bed of solids over a period of 4
			hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
			to yield pale blue colored solid powder.
414.	Copper (II), 4, 4', 4'',4'''-	Saturated strontiun	2 gm bentonite was charged in a pan, which was
	tetrasulfopthalocynine 500	chloride in 5 ml	subsequently set in to rotation. During this procedure
	mg, 500µl ethylene glycol	water	solids were tumbled. Temperature of the rotating pan
l f	and 500 mg sodium silicate in		was raised to 70 °C, under the flow of argon solution A
	5 ml water		was sprayed on the bed of solids over a period of 4
	J III water		hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
		,	to yield pale blue colored solid powder.
415.	Manganese(II), 4, 4', 4'',4'''-	Saturated strontiun	2 gm davisil was charged in a pan, which was
	tetrasulfopthalocynine. 500	chloride in 5 ml	subsequently set in to rotation. During this procedure
l ·	mg, 500µl ethylene glycol	water	solids were tumbled. Temperature of the rotating pan
	and 500 mg sodium silicate in		was raised to 70 °C, under the flow of argon solution A
i i	5 ml water		was sprayed on the bed of solids over a period of 4
•	J III Water		hours followed by spraying solution B. resulting solids
,		,	were further tumbled for 2 hours and dried in vacuum
		•	to yield pale blue colored solid powder.
416		·	
416.	Mangapece(ID A A' A'' A'''-	Saturated harium	2 gm davisil was charged in a pan, which was
1	tetrasulfopthalocynine. 500	nitrate in 5 ml water	subsequently set in to rotation. During this procedure
	mg, 500µl ethylene glycol	maac ii 5 iii waaci	solids were tumbled. Temperature of the rotating pan
1	_		was raised to 70 °C, under the flow of argon solution A
	and 500 mg sodium silicate in		was sprayed on the bed of solids over a period of 4
' 5	5 ml water		hours followed by spraying solution B. resulting solids
			were further tumbled for 2 hours and dried in vacuum
, 1			to yield blue colored solid powder.
			w pleta otao colorea sona pomaca.
417.	fencence(ID 4 4) 49) 49)	Commented beginn	2 gm alumina was charged in a pan, which was
	Manganese(II), 4, 4', 4'',4'''-	Saturated barium	subsequently set in to rotation. During this procedure
1	etrasulfopthalocynine. 500	nitrate in 5 ml water	• •
	ng .		solids were tumbled. Temperature of the rotating pan was raised to 70 °C, under the flow of argon solution A
			was raised to 70 to under the HOW of Argon solition A
	00μl ethylene glycol		• — —
· A	And 500 mg sodium silicate	;	was sprayed on the bed of solids over a period of 4
· A		:	was sprayed on the bed of solids over a period of 4 hours followed by spraying solution B. resulting solids
· A	And 500 mg sodium silicate		was sprayed on the bed of solids over a period of 4

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418.	Manganese(II), 4, 4',	Saturated barium	2 gm alumina was charged in a pan, which
·	4'',4'''-	nitrate in 5 ml	was subsequently set in to rotation. During
	tetrasulfopthalocynine	water	this procedure solids were tumbled.
	500 mg, 500µl ethylene		Temperature of the rotating pan was raised
	glycol and 500 mg		to 70 °C, under the flow of argon solution A
	sodium polyvinyl		was sprayed on the bed of solids over a
	sulfonate in 5 ml water		period of 4 hours followed by spraying
			solution B. resulting solids were further
			tumbled for 2 hours and dried in vacuum to
			yield pale blue colored solid powder.
419.	Iron (III), 4, 4', 4'',4'''-	Saturated	2 gm davisil was charged in a pan, which
	tetrasulfopthalocynine	strontium	was subsequently set in to rotation. During
	oxygen adduct. 500 mg,	chloride in 5 ml	this procedure solids were tumbled.
	500µl ethylene glycol	water	Temperature of the rotating pan was raised
	and 500 mg sodium		to 70 °C, under the flow of argon solution A
	sulfate in 5 ml water		was sprayed on the bed of solids over a
		·	period of 4 hours followed by spraying
			solution B. resulting solids were further
			tumbled for 2 hours and dried in vacuum to
			yield pale blue colored solid powder.
420.	Iron (III), 4, 4', 4'',4'''-	Saturated barium	2 gm davisil was charged in a pan, which
	tetrasulfopthalocynine	nitrate in water 5	was subsequently set in to rotation. During
	oxygen adduct. 500 mg	ml	this procedure solids were tumbled.
	500μl ethylene glycol		Temperature of the rotating pan was raised
•	and 500 mg sodium		to 70 °C, under the flow of argon solution A
	sulfate in 5 ml water		was sprayed on the bed of solids over a
		·	period of 4 hours followed by spraying
			solution B. resulting solids were further
			tumbled for 2 hours and dried in vacuum to
	•		yield pale blue colored solid powder.
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Examples 421 - 429

Catalyst stability in various organic solvents

These examples illustrate the stability of catalysts in liquid phases. Stability of catalyst was assessed in order to establish integrity and resilience of catalyst in liquid phase reactions. Apparatus according to figure 3 was assembled and 5g. catalyst was added in the extraction vessel. 0.5-liter solvent was charged in extraction vessel. Solids in the extractor were agitated and solvent in the round bottomed flask was set to boiling. Solid catalyst was continuously leached for 24 hours. Boiling liquid was brought to room

temperature and analyzed for group IIA metal and transition metal. No leaching of catalytically active material was apparent.

No	Catalyst	Extraction solvent	Observation
421	Catalytic entity: HRhCO(TPPTS)3 (10 ⁻⁶ mols) Additive: TPPTS 6* 10 ⁻⁶ mols/	Water	No leaching detected colour of solid remains unchanged
·	polyvinyl sulfonic acid 100 mg Support: 5 g silica Davisil TM Group IIA metal: barium Method of preparation:	Methanol	No leaching detected colour of solid remains unchanged
	deposition precipitation Colour of catalyst: pale yellow	Acetone	No leaching detected colour of solid remains unchanged
		Thf	No leaching detected colour of solid remains unchanged
		Acetonitrile	No leaching detected colour of solid becomes light orange
·.		DMF	No leaching detected colour of solid becomes light orange
		Chloroform	No leaching detected colour of solid remains unchanged
	·	Tolune	No leaching detected colour of solid remains unchanged
		Hexane	No leaching detected colour of solid remains unchanged
·		Acetic acid	No leaching detected colour of solid remains unchanged

	T		
422	Catalytic entity: Ru(H)(Cl)(TPPTS) ₄ (10 ⁻⁶ mols) Additive: TPPTS 8* 10 ⁻⁶ mols/	Water	No leaching detected colour of solid remains unchanged
	alginic acid 100 mg Support: 5 g silica Davisil ™ Group IIA metal: strontium Method of preparation:	Methanol	No leaching detected colour of solid remains unchanged
	fluidized bed precipitation Colour of catalyst: pale brown	Acetone	No leaching detected colour of solid remains unchanged
		Thf	No leaching detected colour of solid remains unchanged
		Acetonitrile	No leaching detected colour of solid turns slightly dark
		DMF	No leaching detected colour of solid changes slightly dark
		Chloroform	No leaching detected colour of solid remains unchanged
		Toluene	No leaching detected colour of solid remains unchanged
		Hexane	No leaching detected colour of solid remains unchanged
·		Acetic acid	No leaching detected colour of solid remains unchanged

			
423	Catalytic entity: palladium acetate bypyridine disulfonated 10 ⁻⁶ mols	Water	No leaching detected colour of solid remains unchanged
	Additive: bypyridine disulfonated 10 ⁻⁵ mols/ polyacrylic acid acid 100 mg Support: 5 g charcoal	Methanol	No leaching detected colour of solid remains unchanged
	Group IIA metal: strontium Method of preparation: fluidized bed precipitation Colour of catalyst: pale orange	Acetone	No leaching detected colour of solid remains unchanged.
		THF	No leaching detected colour of solid remains unchanged
		Acetonitrile	No leaching detected colour of solid remains unchanged
		DMF	No leaching detected colour of solid remains unchanged
		Chloroform	No leaching detected colour of solid remains unchanged
		Toluene	No leaching detected colour of solid remains unchanged
		Hexane	No leaching detected colour of solid remains unchanged
		Acetic acid	No leaching detected colour of solid remains unchanged

			· ·
424	Catalytic entity:	Water	No leaching detected
•	cobalt(II)4,4',4'',4''',-		colour of solid remains
	tetrasulfopthalocynine 10 ⁻⁶ mols		unchanged
	additive: sodium phosphate 100		
	mg	Methanol	No leaching detected
	Support: 5 g gamma alumina		colour of solid remains
	Group IIA metal: barium		unchanged
	Method of preparation: coating		
	pan precipitation	Acetone	No leaching detected
	Colour of catalyst: pale blue		colour of solid remains
			unchanged
	•	THF	No leaching detected
			colour of solid remains
			unchanged
			anonangou .
		Acetonitrile	No leaching detected
		·	colour of solid remains
			unchanged
			unonungou
*		DMF	No leaching detected
		DIVII	colour of solid remains
			unchanged
	l ·		unchanged
		Chloroform	No leaching detected
•	·	Cinorotoriii	colour of solid remains
			•
			unchanged
		Tal	No looshing days and
•	·	Toluene	No leaching detected
			colour of solid remains
•	.		unchanged
		Hexane	No leaching detected
			colour of solid remains
			unchanged
		Acetic acid	No leaching detected
	·		colour of solid remains
			unchanged

			
425	Catalytic entity: sulfonated quaternary ammonium	Water	No leaching detected
	hydroxide benzyl triphenyl amine 10 ⁻⁴ mols	Methanol	No leaching detected
	additive: carboxymethyl cellulose 100 mg	Acetone	No leaching detected
	Support: 5 g activated charcoal Group IIA metal: barium	тнг	No leaching detected
	Method of preparation: coprecipitation	Chloroform	No leaching detected
		Toluene	No leaching detected
		Hexane	No leaching detected
426	Catalytic entity: cobalt(II) N,N'- ethylene bis(salicyldiamine 5- sodium sulfonate) 10 ⁻⁴ mols additive: sodium sulfate 200 mg	Water	No leaching detected colour of solid remains unchanged
·	Support: 5 g asbestos Group IIA metal: barium Method of preparation: coating pan	Methanol	No leaching detected colour of solid remains unchanged
	Color of the catalyst: gray brown	Acetone	No leaching detected colour of solid remains unchanged
		THF	No leaching detected colour of solid remains unchanged
		Acetonitrile	No leaching detected colour of solid remains unchanged
		DMF	No leaching detected colour of solid remains unchanged

•• .		Chloroform	No leaching detected colour of solid remains
			unchanged
		Toluene	No leaching detected
			colour of solid remains
			unchanged
		TT	No looking detected
		Hexane	No leaching detected colour of solid remains
		•	unchanged
			unchanged
		Acetic acid	No leaching detected
			colour of solid remains
			unchanged
427	Catalytic entity:	Water	No leaching detected
•	NiCl₂(TPPTS)₂10 ⁻⁴ mols	Methanol	No leaching detected
	additive: tppts 20* 10 ⁻⁴ mols	Acetone	No leaching detected
	Support: 5 g activated charcoal	THF	No leaching detected
	Group IIA metal: strontium	Acetonitrile	No leaching detected
	Method of preparation:	DMF	No leaching detected
	fluidized bed precipitation	Chloroform	No leaching detected
		Tolune	No leaching detected
		Hexane	No leaching detected
		Acetic acid	No leaching detected
428	Catalytic entity: Rh *ClO ₄ * (S,S)		No leaching detected
	chiraphos tetra sulfonated 10 ⁻⁶	Water	colour of solid remains
	mols		unchanged
	additive: s,s chiraphos 20* 10 ⁻⁶	76.1	No leaching detected
	mols	Methanol	colour of solid remains
	Support: 5 g silica Davisil TM		unchanged
	Group IIA metal: barium Method of preparation: coating	Acetone	No leaching detected
	pan precipitation	Activité	colour of solid remains
	Color of the catalyst: pale		unchanged
	yellow	 	No leaching detected
		Thf	colour of solid remains
			unchanged

		Acetonitrile	No leaching detected
1		Accionatio	colour of solid remains
			unchanged
	·	DMF	No leaching detected
			colour of solid remains
			unchanged
		Chloroform	No leaching detected
			colour of solid remains
			unchanged
		Tolune	No leaching detected
	·		colour of solid remains
			unchanged
		Hexane	No leaching detected
	·		colour of solid remains
			unchanged
		Acetic acid	No leaching detected
			colour of solid remains
			unchanged
429	Catalytic entity: MoO ₄ -2 10 -4	Water	No leaching detected
	mols		colour of solid remains
	additive: sodium phosphate 500		unchanged
	mg	Methanol	No leaching detected
	Support: 5 g calcium silicate		colour of solid remains
	Group IIA metal: calcium	·	unchanged
	Method of preparation:	Acetone	No leaching detected
	coprecipitation	,	colour of solid remains
		۲.	unchanged
		Thf	No leaching detected
			colour of solid remains
			unchanged
		Acetonitrile	No leaching detected
, ,			colour of solid remains
			unchanged
.		DMF	No leaching detected
		DIMI.	colour of solid remains
			unchanged
'	•		dichanged
		Chloroform	No leaching detected
	_		colour of solid remains
]		•	unchanged
			

·	Tolune	No leaching detected colour of solid remains unchanged
	Hexane	No leaching detected colour of solid remains unchanged
	Acetic acid	No leaching detected colour of solid remains unchanged

Example 430

Hydroformylation reaction as a probe

This example illustrates the applicability of the catalytic formulation to liquid phase reaction wherein two gases react with substrate in liquid phase. This example also illustrates how solid catalyst can be employed to catalyze reaction and a preferred method to recover and recycle it.

Catalyst specification

Catalytic entity	HRhCO(TPPTS)3 (10.8 * 10 ⁻⁶ mols)	
Additive	TPPTS 6* 10 ⁻⁶ mols/ polyvinyl sulfonic	
	acid 100 mg	
Support:	5 g silica Davisil TM	
Group IIA metal	barium	
Method of preparation	deposition precipitation	
Colour of catalyst	pale yellow	
Metal content in solid	10.8 * 10 ⁻⁶ mols of rhodium.	

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Reaction procedure: Under a argon atmosphere the micro-reactor was charged with 1 g. of catalyst and 25 ml of octene the reactor was flushed with H_2 / CO mixture and reactor was heated to 75^0 C and pressurized with H_2 / CO mixture (1:1by mole) to 600 psi and maintained at this temperature. Liquid suspension was stirred at 900 rpm. Reaction was continued for 240 min. analysis of the products confirmed conversion of olefins to aldehydes.

Conversion 14 * 10⁻³ mols of octene

Turn over frequencies at 60 min were 894 hour⁻¹

Turnover number after 240 min was 1296.29 mols. mol⁻¹ of catalyst

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n/i ratio after 240 min was 2.7 (wherein n is linear aldehyde and I is iso aldehyde)
Color of recovered catalyst: light brown

The catalyst was recovered by centrifugation and repeatedly washing reactor with tolune under nitrogen atmosphere. Solid catalyst was dried under vacuum. Which was recycled to perform reaction as described earlier to obtain equivalent activity and selectivity. The color of catalyst was light brown

Examples 431

Need of the support

These comparative examples illustrate the need of solid support in the catalytic formulation and effect of loading of catalytically active solid material on solid support to decide a protocol for optimum loading.

Hydroformylation of hexene

Catalyst specifications:

Catalytic entity	HRhCO(TPPTS)3 (10 ⁻⁶ mols)	
Additive	TPPTS 6* 10 ⁻⁶ mols/ polyvinyl sulfonic	
	acid 100 mg	
Support:	5 g silica Davisil TM	
Group IIA metal	barium	
Method of preparation	deposition precipitation	
Colour of catalyst	pale yellow	

Procedure:

Under a argon atmosphere the microreactor was charged with 2 g of catalyst and 0.5 g (5.2 * 10^{-4} mols) of hexene in 20 ml tolune the reactor was flushed with H_2 / CO mixture and reactor was heated to 75 0 C and pressurized with H_2 / CO mixture (1:1by mole) and maintained at this temperature. Liquid suspension was stirred with external magnetic agitation. Reaction was continued for 24 hours. Analysis of the products confirmed conversion of olefins to aldehydes. 89 % Olefin was converted to aldehydes with n/I selectivity of 1.80.

Under identical conditions 4 * 10⁻⁷ mols of barium salt of rhodium catalyst failed to promote any reaction.

Example 432

25 Effect of the added ligand

These comparative examples illustrate the need of additional ligand in catalytic formulation. These examples also illustrate solid support in the catalytic formulation and

effect of loading of catalytically active solid material on solid support to decide optimum loading.

Catalyst preparation catalyst of varying specifications were prepared by following method:

Hydroformylation of hexene

Preparation of catalyst:

Catalyst specifications:

	Catalyst A	Catalyst B	Catalyst C
Catalytic entity	HRhCO(TPPTS)3	HRhCO(TPPTS)3	HRhCO(TPPTS)3
·	(10 ⁻⁶ mols)	(10 ⁻⁶ mols)	(10 ⁻⁶ mols)
		·	
Additive	polyvinyl sulfonic	TPPTS 6* 10 ⁻⁶ mols/	TPPTS 12* 10 ⁻⁶
·	acid 100 mg	polyvinyl sulfonic	mols/ polyvinyl
		acid 100 mg	sulfonic acid 100 mg
Support:	5 g silica Davisil ™	5 g silica Davisil TM	5 g silica Davisil TM
Group IIA metal	barium	barium	barium
Method of	deposition	deposition	deposition
preparation	precipitation	precipitation	precipitation
Colour of catalyst	pale yellow	pale yellow	pale yellow

10 Procedure:

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Under a argon atmosphere the microreactor was charged with 2 g of catalyst and $0.5 \text{ g} (5.2 * 10^{-4} \text{ mols})$ of hexene in 20 ml tolune the reactor was flushed with H₂/CO mixture and reactor was heated to 75 0 C and pressurized with H₂/CO mixture (1:1by mole) and maintained at this temperature. Liquid suspension was stirred with external magnetic agitation. Reaction was continued for 24 hours.

Analysis of the products confirmed conversion of olefins to aldehydes. About 89 % Olefin was converted to aldehydes with n/I selectivity of 1.80.

Example 433

Effect of added water-Hydroformylation of hexene

Catalyst specifications:

Catalytic entity	HRhCO(TPPTS)3 (10 ⁻⁶ mols)
Additive	TPPTS 6* 10 ⁻⁶ mols/ polyvinyl sulfonic
	acid 100 mg
Support:	5 g silica Davisil TM
Group IIA metal	Barium
Method of preparation	deposition precipitation
Colour of catalyst	pale yellow
Moisture content	2%

Procedure:

Under a argon atmosphere the microreactor was charged with 2 g of catalyst and 0.5 g (5.2 * 10^{-4} mols) of hexene in 20 ml tolune the reactor was flushed with H₂/ CO mixture and reactor was heated to 75° C and pressurized with H₂/ CO mixture (1:1by mole) and maintained at this temperature. Liquid suspension was stirred with external magnetic agitation. Reaction was continued for 24 hours. Analysis of the products confirmed conversion of olefins to aldehydes. 89% Olefin was converted to aldehydes with n/I selectivity of 1.80.

Under identical conditions in reaction charge 1g water was added and after 24 hours only 5% conversion was obtained with similar n/I ratio.

Example 434

Continuous fixed bed experiment

15 Catalyst specifications:

Catalytic entity	HRhCO(TPPTS)3 (10 ⁻⁶ mols)
Additive	TPPTS 6* 10 ⁻⁶ mols/ polyvinyl sulfonic
	acid 100 mg
Support:	5 g silica Davisil TM
Group IIA metal	barium
Method of preparation	deposition precipitation
Colour of catalyst	pale yellow

Procedure: Accordingly the crucial evaluation indicating life of the catalyst, its stability and the durability was performed in a tubular fixed bed reactor by subjecting catalyst to

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hydroformylation in tubular trical bed reactor (ϕ ½") at 80 0 C and 300 psi H₂/CO (1:1) using 5 g. of catalyst. 5 % decene in toluene was pumped continuously at the feed rate of 10 ml/hr conversion levels were 20 %(fluctuating by +-2.2 %) for aldehydes (n/i 2.1) after attaining steady state (5 hours). The reaction was further continued for 76 hr without loss of activity. Reaction was arrested by discontinuing the liquid feed and water was pumped for 1 hr. thereafter reactant feed was resumed. Initially there was no conversion, which was steadily resumed to earlier levels over the period of 10 hr. This observation was attributed to formation of water film on the catalyst surface, which physically retards contact of decene with catalyst surface. Moreover water does not wash out complex catalyst, which provides conclusive proof that reaction occurs in the solid state.

Example 435

Hydroformylation of hexene

Catalyst specifications:

Catalytic entity	HRhCO(TPPTS)3 (10 ⁻⁶ mols)	
Additive	TPPTS 6* 10 ⁻⁶ mols/ polyvinyl sulfonic	
	acid 100 mg	
Support:	5 g silica Davisil TM	
Group IIA metal	barium	
Method of preparation	deposition precipitation	
Color of catalyst	pale yellow	

Procedure:

Under a argon atmosphere the microreactor was charged with 200 mg of catalyst and 0.05 g (5.2 * 10⁻⁴ mols) of hexene in 2 ml toluene the reactor was flushed with H₂/CO mixture and reactor was heated to 75 °C and pressurized with H₂/CO mixture (1:1by mole) and maintained at this temperature. Liquid suspension was stirred with external magnetic agitation. Reaction was continued for 24 hours. Analysis of the products confirmed conversion of olefins to aldehydes. 89 % Olefin was converted to aldehydes with n/I selectivity of 1.91 the catalyst was recovered by washing reactor several times with toluene and catalyst was recovered by centrifugation, washing repeatedly with toluene and diethyl ether. Catalyst was dried under vacuum and recycled to obtain equivalent activity.

Example 436

25 Hydroformylation of styrene

Preparation of catalysts

Catalyst specifications:

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Catalytic entity	HRhCO(TPPTS)3 (10 ⁻⁶ mols)
Additive	TPPTS 6* 10 ⁻⁶ mols/ polyvinyl sulfonic acid 100
	mg
Support:	5 g silica Davisil ™
Group IIA metal	barium
Method of preparation	deposition precipitation
Color of catalyst	pale yellow

Under a argon atmosphere the microreactor was charged with 200 mg of catalyst and 0.05 g (4.8 * 10 ⁻⁴) of styrene in 2 ml toluene the reactor was flushed with H₂/ CO mixture and reactor was heated to 75 °C and pressurized with H₂/ CO mixture (1:1by mole) and maintained at this temperature. Liquid suspension was stirred with external magnetic agitation. Reaction was continued for 24 hours. Analysis of the products confirmed conversion of olefins to aldehydes. 91 % Olefin was converted to aldehydes with n/I ratio of 0.449. The catalyst was recovered by washing reactor several times with toluene and catalyst was recovered by centrifugation, washing repeatedly with toluene and diethyl ether. Catalyst was dried under vacuum and recycled to obtain equivalent activity.

Example 437

Hydroformylation of cyclohexene

Preparation of catalysts

Catalyst specifications:

Catalytic entity	HRhCO(TPPTS)3 (10 ⁻⁶ mols)
Additive	TPPTS 6* 10 ⁻⁶ mols/ polyvinyl sulfonic acid 100
	mg
Support:	5 g silica Davisil TM
Group IIA metal	barium
Method of preparation	deposition precipitation
Color of catalyst	pale yellow

Under a argon atmosphere the microreactor was charged with 200 mg of catalyst and 0.05 g (6.1 * 10^{-4}) of cyclohexene in 2 ml toluene the reactor was flushed with H₂/ CO mixture and reactor was heated to 75 0 C and pressurized with H₂/ CO mixture (1:1by mole) and maintained at this temperature. Liquid suspension was stirred with external magnetic

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agitation. Reaction was continued for 24 hours. Analysis of the products confirmed conversion of olefins to aldehydes.47 % Olefin was converted to aldehydes. The catalyst was recovered by washing reactor several times with toluene and catalyst was recovered by centrifugation, washing repeatedly with toluene and diethyl ether. Catalyst was dried under vacuum and recycled to obtain equivalent activity.

Example 438

Hydroformylation of allyl alcohol

Catalyst specifications:

Catalytic entity	HRhCO(TPPTS)3 (10 ⁻⁶ mols)
Additive	TPPTS 6* 10 ⁻⁶ mols/ polyvinyl sulfonic
	acid 100 mg
Support:	5 g silica Davisil TM
Group IIA metal	barium
Method of preparation	deposition precipitation
Color of catalyst	pale yellow

Procedure: Under a argon atmosphere the microreactor was charged with 200 mg of catalyst and 0.05 g (8.7 * 10⁻⁴ mols) of allyl alcohol in 2 ml water the reactor was flushed with H₂/ CO mixture and reactor was heated to 75 °C and pressurized with H₂/ CO mixture (1:1by mole) and maintained at this temperature. Liquid suspension was stirred with external magnetic agitation. Reaction was continued for 24 hours. Analysis of the products confirmed conversion of olefins to aldehydes. 20 % Olefin was converted to aldehydes with n/i ratio of 1. The catalyst was recovered by washing reactor several times with toluene and catalyst was recovered by centrifugation, washing repeatedly with toluene and diethyl ether. Catalyst was dried under vacuum and recycled to obtain equivalent activity.

Example 439

Hydroformylation of hexene

Catalyst specifications:

Catalytic entity	HRhCO(N(Ph _m SO ₃ ⁻) ₃) ₃ (10 ⁻⁶ mols)
Additive	N(Ph _m SO ₃ ⁻) ₃ 6* 10 ⁻⁶ mols/ polyvinyl
	sulfonic acid 100 mg

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Support	5 g silica Davisil TM
Group IIA metal salt	barium
Method of preparation	deposition precipitation
Color of catalyst	pale yellow

Under a argon atmosphere the microreactor was charged with 200 mg of catalyst and 0.5 g (5.9 *10⁻⁴, mol) of hexene in 2 ml toluene the reactor was flushed with H₂/ CO mixture and reactor was heated to 75 °C and pressurized with H₂/ CO mixture (1:1by mole) and maintained at this temperature. Liquid suspension was stirred with external magnetic agitation. Reaction was continued for 24 hours. Analysis of the products confirmed conversion of olefins to aldehydes. 68 % Olefin was converted to aldehydes with n/I selectivity of 1.78 the catalyst was recovered by washing reactor several times with toluene and catalyst was recovered by centrifugation, washing repeatedly with toluene and diethyl ether. Catalyst was dried under vacuum and recycled to obtain equivalent activity.

Example 440

Hydroformylation of hexene

Catalyst specifications:

Catalytic entity	HRhCO(BISBIS) (10 ⁻⁶ mols)
Additive	BISBIS 6* 10 ⁻⁶ mols/ polyvinyl sulfonic
	acid 100 mg
Support	5 g silica Davisil TM
Group IIA metal salt	barium
Method of preparation	deposition precipitation
Color of catalyst	pale yellow

Under a argon atmosphere the microreactor was charged with 200 mg of catalyst and 0.05 g (5.9 * 10^{-4} , mol) of hexene in 2 ml toluene the reactor was flushed with H₂/ CO mixture and reactor was heated to 75°C and pressurized with H₂/ CO mixture (1:1by mole) and maintained at this temperature. Liquid suspension was stirred with external magnetic

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agitation. Reaction was continued for 24 hours. Analysis of the products confirmed conversion of olefins to aldehydes. 78 % Olefin was converted to aldehydes with n/I selectivity of 17.88. The catalyst was recovered by washing reactor several times with toluene and catalyst was recovered by centrifugation, washing repeatedly with toluene and diethyl ether. Catalyst was dried under vacuum and recycled to obtain equivalent activity.

Example 441

Hydroformylation of hexene

Catalyst specifications:

Catalytic entity	HrhCO(succindiphos) (10 ⁻⁶ mols)
Additive	succindiphos 6* 10 ⁻⁶ mols/ polyvinyl
	sulfonic acid 100 mg
Support	5 g silica Davisil TM
Group IIA metal salt	barium
Method of preparation	deposition precipitation
Color of catalyst	pale yellow

Under a argon atmosphere the microreactor was charged with 200 mg of catalyst and 0.05 g (5.9 *10⁻⁴, mol) of hexene in 2 ml toluene the reactor was flushed with H₂/CO mixture and reactor was heated to 75⁰C and pressurized with H₂/CO mixture (1:1by mole) and maintained at this temperature. Liquid suspension was stirred with external magnetic agitation. Reaction was continued for 24 hours. Analysis of the products confirmed conversion of olefins to aldehydes. 80 % Olefin was converted to aldehydes with n/I selectivity of 0.6. The catalyst was recovered by washing reactor several times with toluene and catalyst was recovered by centrifugation, washing repeatedly with toluene and diethyl ether. Catalyst was dried under vacuum and recycled to obtain equivalent activity.

Example 442

20 Hydroformylation of hexene

Catalyst specifications:

Catalytic entity	HRhCO(bypyds) (10 ⁻⁶ mols)
Additive	bypyds 6* 10 ⁻⁶ mols/ polyvinyl sulfonic
	acid 100 mg

Support	5 g silica Davisil TM
Group IIA metal salt	barium
Method of preparation	deposition precipitation
Color of catalyst	pale yellow

Under a argon atmosphere the microreactor was charged with 200 mg of catalyst and 0.05 g (5.9 *10⁻⁴, mol) of hexene in 2 ml toluene the reactor was flushed with H₂/ CO mixture and reactor was heated to 75 °C and pressurized with H₂/ CO mixture (1:1by mole) and maintained at this temperature. Liquid suspension was stirred with external magnetic agitation. Reaction was continued for 24 hours. Analysis of the products confirmed conversion of olefins to aldehydes. 80 % Olefin was converted to aldehydes with n/I selectivity of 0.92. The catalyst was recovered by washing reactor several times with toluene and catalyst was recovered by centrifugation, washing repeatedly with toluene and diethyl ether. Catalyst was dried under vacuum and recycled to obtain equivalent activity.

Example 443

Cobalt catalyzed hydroformylation

Catalyst specifications:

Catalytic entity	$(\text{Co(Ac)}_2/(\text{P(Ph_mSO}_3)_3)_3 \ (10^{-6} \text{ mols})$
Additive	P(Ph _m SO ₃) ₃ 6* 10 ⁻⁶ mols/ polyvinyl
Support	sulfonic acid 100 mg 5 g silica Davisil TM
Group IIA metal salt	barium
Method of preparation	deposition precipitation
Color of catalyst	pale yellow

Under a argon atmosphere the microreactor was charged with 200 mg of catalyst and 0.05 g (5.9 * 10^{-4} , mol) of hexene in 2 ml toluene the reactor was flushed with H₂/CO mixture and reactor was heated to 75°C and pressurized with H₂/CO mixture (1:1by mole) and

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maintained at this temperature. Liquid suspension was stirred with external magnetic agitation. Reaction was continued for 24 hours. Analysis of the products confirmed conversion of olefins to aldehydes. 30 % Olefin was converted to aldehydes with n/i of 1.38. The catalyst was recovered by washing reactor several times with toluene and catalyst was recovered by centrifugation, washing repeatedly with toluene and diethyl ether. Catalyst was dried under vacuum and recycled to obtain equivalent activity.

Example 444

Platinum catalyzed hydroformylation

Preparation of catalysts: heterogenized platinum chloride phosphine complex was boiled with dichloromethane and stannous chloride and subsequently extracted with dichloromethane

Catalyst specifications:

Catalytic entity	$SnCl_2PtCl_2(P(Ph_mSO_3)_3)_2 (10^{-6} mols)$
Additive	N(Ph _m SO ₃ ⁻) ₃ 6* 10 ⁻⁶ mols/ polyvinyl
	sulfonic acid 100 mg
Support	5 g silica Davisil TM
Group IIA metal salt	barium
Method of preparation	deposition precipitation
Color of catalyst	pale yellow

Procedure: Under a argon atmosphere the microreactor was charged with 200 mg of catalyst and 0.05 g (5.9 *10⁻⁴, mol) of hexene in 2 ml toluene the reactor was flushed with H₂/ CO mixture and reactor was heated to 75 °C and pressurized with H₂/ CO mixture (1:1by mole) and maintained at this temperature. Liquid suspension was stirred with external magnetic agitation. Reaction was continued for 24 hours. Analysis of the products confirmed conversion of olefins to aldehydes. 57 % Olefin was converted to aldehydes with n/I selectivity of 10.47. The catalyst was recovered by washing reactor several times with toluene and catalyst was recovered by centrifugation, washing repeatedly with toluene and diethyl ether. Catalyst was dried under vacuum and recycled to obtain equivalent activity.

Example 445

Carbonylation of styrene

Catalyst specifications:

Catalytic entity	$Pd(Ac)_2(P(Ph_mSO_3^-)_2(PySO_3^-))$ (10 ⁻⁶
Additive	mols) (P(Ph _m SO ₃) ₂ (PySO ₃) 6* 10 ⁻⁶ mols/
	polyvinyl sulfonic acid 100 mg
Support	5 g charcoal
Group IIA metal salt	barium
Method of preparation	deposition precipitation
Color of catalyst	Black
Metal content	

Procedure: 15 ml reactor with magnetic stirrer bar was charged with 200 mg catalyst and 500 mg (4.90 10⁻³ mmol) styrene and 5 mg p toluene sulfonic, acid 25 mg N, N, dimethyl aniline and 10 ml methanol. Micro reactor was flushed with argon and pressurized with carbon monoxide 800 psi and mixture was stirred at 70 °C for 24 hours. Reaction mixture was analyzed by gas chromatograph. 87 % phenyl acetylene was carbonylated with 99 % selectivity for methyl 2phenyl propionate. Reaction mixture was centrifuged to recover solid catalyst, which was subsequently washed with methanol repeatedly. Catalyst was further washed with diethyl ether and dried under vacuum.

Example 446

15 Carbonylation of styryl alcohol

Preparation of catalysts

Catalyst specifications:

$Pd(Ac)_2(P(Ph_mSO_3^-)_2(PySO_3^-))$ (10 ⁻⁶ mols)
(P(Ph _m SO ₃) ₂ (PySO ₃) 6* 10 ⁻⁶ mols/ polyvinyl sulfonic acid 100 mg
5 g charcoal
barium
deposition precipitation
Black

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Procedure: 15 ml reactor with magnetic stirrer bar was charged with 200 mg catalyst and 500 mg (8.9 * 10⁻³ mmol) styryl alcohol and 5 mg p toluene sulfonic, acid and 10 ml methanol. Micro reactor was flushed with argon and pressurized with carbon monoxide 800 psi and mixture was stirred at 100 °C for 24 hours. Reaction mixture was analyzed by gas chromatograph. 52 % phenyl acetylene was carbonylated with 91 % selectivity for methyl 2 phenyl propionate over 2-phenyl propionate. Reaction mixture was centrifuged to recover solid catalyst, which was subsequently washed with methanol repeatedly. Catalyst was further washed with diethyl ether and dried under vacuum.

Example 447

10 Carbonylation of phenyl acetylene

Catalyst specifications:

Catalytic entity	$\mathbf{Pd(Ac)_2}(P(Ph_mSO_3)_2(PySO_3)) (10^{-6} \text{ mols})$
Additive	(P(Ph _m SO ₃) ₂ (PySO ₃) 6* 10 ⁻⁶ mols/ polyvinyl sulfonic acid 100 mg
Support	5 g charcoal
Group IIA metal salt	barium
Method of preparation	deposition precipitation
Color of catalyst	Black

Procedure: 15 ml reactor with magnetic stirrer bar was charged with 200 mg catalyst and 500 mg (4.90 10⁻³ mmol) phenyl acetylene and 5 mg p toluene sulfonic, acid 25 mg N, N, dimethyl aniline and 10 ml methanol. Micro reactor was flushed with argon and pressurized with carbon monoxide 100 psi and mixture was stirred at 90 °C for 12 hours. Reaction mixture was analyzed by gas chromatograph. 80 % phenyl acetylene was carbonylated with 96 % selectivity for methyl dehydroatropate. Reaction mixture was centrifuged to recover solid catalyst, which was subsequently washed with methanol repeatedly. Catalyst was further washed with diethyl ether and dried under vacuum.

Example 448

Hydrogenation of styrene

Catalyst specifications:

Catalytic entity		RhClCOD(TPPTS) ₃ 10 ⁻⁵ mols
Additive	•	10 ⁻⁵ mols of tppts/ 500 mg sodium carboxy
		methyl cellulose

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Support	kesilghur
Group IIA metal salt	Strontium chloride saturated solution
Method of preparation	Deposition precipitation
Color of catalyst	Pale yellow

Procedure: 15 ml reactor with magnetic stirrer bar was charged with 200 mg catalyst and 500 mg (4.8 * 10⁻³ mmol) styrene in 10 ml ethanol. Micro reactor was flushed with argon and pressurized with hydrogen 500 psi and mixture was stirred at 90 °C for 12 hours. Reaction mixture was analyzed by gas chromatograph. 98 % styrene was hydrogenated to ethyl benzene. Reaction mixture was centrifuged to recover solid catalyst, which was subsequently washed with methanol repeatedly. Catalyst was further washed with diethyl ether and dried under vacuum. Catalyst was recycled to obtain equivalent activity

Example 449

Hydrogenation of methyl cinnamate

Catalyst specifications:

Catalytic entity	RhClCOD(TPPTS) ₃ 10 ⁻⁵ mols
Additive	10 ⁻⁵ mols of tppts/ 500 mg sodium carboxy
	methyl cellulose
Support	kesilghur
Group IIA metal salt	Strontium chloride saturated solution
Method of preparation	Deposition precipitation
Color of catalyst	Pale yellow

Procedure: 15 ml reactor with magnetic stirrer bar was charged with 200 mg catalyst and 500 mg (3.08 * 10⁻³ mmol) methyl cinnamate in 10 ml methanol. Micro reactor was flushed with argon and pressurized with hydrogen 1000 psi and mixture was stirred at 50⁰C for 12 hours. Reaction mixture was analyzed by gas chromatograph. 80% methyl cinnamate was hydrogenated to methyl 3 phenyl propionate. Reaction mixture was centrifuged to recover solid catalyst, which was subsequently washed with methanol repeatedly. Catalyst was further washed with diethyl ether and dried under vacuum. Catalyst was recycled to obtain equivalent activity.

Example 450

Hydrogenation of cinnamonitrile Catalyst specifications:

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Catalytic entity	RhClCOD(TPPTS) ₃ 10 ⁻⁵ mols
Additive	10 ⁻⁵ mols of tppts/ 500 mg sodium carboxy methyl cellulose
Support	Kesilghur
Group IIA metal salt	Strontium chloride saturated solution
Method of preparation	Deposition precipitation
Color of catalyst	Pale yellow

Procedure: 15 ml reactor with magnetic stirrer bar was charged with 200 mg catalyst and 500 mg (3.87 * 10⁻³ mmol) cinnamonitrile in 10 ml methanol. Micro reactor was flushed with argon and pressurized with hydrogen 500 psi and mixture was stirred at 50 °C for 12 hours. Reaction mixture was analyzed by gas chromatograph. 79 % cinnamonitrile was hydrogenated with 60 % selectivity for 3-phenyl propionitrile. Reaction mixture was centrifuged to recover solid catalyst, which was subsequently washed with methanol repeatedly. Catalyst was further washed with diethyl ether and dried under vacuum. Catalyst was recycled to obtain equivalent activity.

Example 451

Hydrogenation of dehydronaproxen

Catalyst specifications:

Catalytic entity	BINAPts RuCl ₂ 10 ⁻⁶ mols
Additive	BINAPts 10 ⁻⁶ mols/ 500 mg sodium phosphate
Support	Silica davisil 1g
Group IIA metal salt	Saturated solution of barium nitrate
Method of preparation	Precipitation in coating pan
Color of catalyst	Pale yellow

Procedure: 15 ml reactor with magnetic stirrer bar was charged with 200 mg catalyst and 128 mg (1.26 * 10⁻³ mol) dehydro naproxen and 128 mg (1.26 * 10⁻³ mol) triethyl amine and 10 ml tolune: methanol (3:2 v/v). Micro reactor was flushed with argon and pressurized with hydrogen 100 bar and mixture was stirred at 0°C for 48 hours. Reaction mixture was centrifuged to recover solid catalyst, which was subsequently washed with methanol repeatedly. All washings and reaction mixture were combined and dried in

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vacuum. Solid thus obtained was thus dissolved in dichloromethane and washed with dilute HCl followed by water. Dichloromethane was evaporated to obtain naproxen.

Analysis of products: products were analyzed by HPLC with WHELK-O column (produced by Merck) yield of naproxen was 98 % and 92% e.e

Second recycle 98% and e.e. 94%

Example 452

Hydrogenation of heptaldehyde

Catalyst specifications:

Catalytic entity	Ru(H)(Cl)(TPPTS) ₃ (10 ⁻⁶ mols)
Additive	TPPTS (6 * 10 ⁻⁶ mols)/ 500 mg sodium alginate
Support	Titania 5 g
Group IIA metal salt	Strontium chloride saturated solution
Method of preparation	coprecipitation
Color of catalyst	Pale yellow

Procedure: 100 mg of the catalyst was charged in the microreactor to this 50 mg (4.38 * 10^{-4} mol) heptaldehyde was added as solution in 2 ml toluene. Microreactor was flushed with argon and heated to 90 °C. Magnetic agitation was started. After attaining temperature reactor was pressurized with 500-psi hydrogen. The reactor was maintained under these conditions for 24 hours. Reaction was stopped by cooling reactor to 0 °C and depressurizing. Reactor was opened and liquid was analyzed by gas chromatograph. 99 % heptaldehyde was converted.

Catalyst was recovered by washing reactor with several portions of toluene combined fractions were centrifuged to recover catalyst. Recovered catalyst was washed with water, methanol and diethyl ether and catalyst was dried under vacuum. This catalyst was recycled to obtain equivalent activity and selectivity. Catalyst was recycled to obtain equivalent conversion and selectivity.

Example 453

Hydrogenation of cinnamaldehyde

Catalyst specifications:

Catalytic entity	Ru(H) ₂ (TPPTS) ₄ (10 ⁻⁶ mols)
Additive	TPPTS (6 * 10 ⁻⁶ mols)/ 500 mg sodium alginate
Support	Titania 5 g
Group IIA metal salt	Strontium chloride saturated solution
Method of preparation	coprecipitation
Color of catalyst	Pale yellow

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Procedure: 100 mg of the catalyst was charged in the microreactor to this 500 mg (3.78 * 10⁻³ mol) cinnamaldehyde was added as solution in 2 ml tetrahydrofuran. Microreactor was flushed with argon and heated to 90 °C. Magnetic agitation was started. After attaining temperature reactor was pressurized with 500-psi hydrogen. The reactor was maintained under these conditions for 24 hours. Reaction was stopped by cooling reactor to 0 °C and depressurizing. Reactor was opened and liquid was analyzed by gas chromatograph. 88 % cinnamaldehyde was converted. Selectivity for cinnamyl alcohol was 73 %.

Catalyst was recovered by washing reactor with several portions of tetrahydrofuran combined fractions were centrifuged to recover catalyst. Recovered catalyst was washed with water, methanol and diethyl ether and catalyst was dried under vacuum. This catalyst was recycled to obtain equivalent activity and selectivity. Catalyst was recycled to obtain equivalent conversion and selectivity.

Example 454

Hydrogenation of ethyl acetoacetate

Catalyst specifications:

Catalytic entity	BINAPts RuCl ₂ 10 ⁻⁶ mols
Additive	BINAPts 10 ⁻⁶ mols/ 500 mg sodium phosphate
Support	Silica davisil 1g
Group IIA metal salt	Saturated solution of barium nitrate
Method of preparation	Precipitation in coating pan
Color of catalyst	Pale yellow

Procedure: Procedure: 1g of the catalyst was charged in the microreactor to this 5g ethyl 3 oxobutanoate in 15 ml methanol. Resulting suspension was charged in microreactor, which was flushed with argon and hydrogen. Temperature of the reactor was raised to 90 °C and reactor was filled with hydrogen 150 psi. The reactor was maintained under these conditions for 24 hours. Reaction was stopped by cooling reactor to 0 °C and depressurizing. Reactor was opened and liquid was analyzed by gas chromatograph. Ethyl 3 oxobutanoate was quantitatively converted to corresponding alcohol. Catalyst was recovered by washing reactor with several portions of methanol. Combined fractions were fractionated to recover catalyst. Recovered catalyst was washed with water, methanol and diethyl ether and catalyst was dried under vacuum. This catalyst was recycled to obtain

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equivalent activity and selectivity. Catalyst was recycled to obtain equivalent conversion and selectivity.

Optical purity of product by fresh catalyst is 92 % and recycled catalyst is 94 %.

Example 455

Hydrogenation of benzyledene acetone

Catalyst specifications:

Catalytic entity	Rh(ClO ₄)(CODI)(TPPTS) ₃ (10 ⁻⁶ mols)
Additive	TPPTS (6 * 10 ⁻⁶ mols)/ 500 mg sodium alginate
Support	zirconia 5 g
Group IIA metal salt	Strontium chloride saturated solution
Method of preparation	coprecipitation
Color of catalyst	Pale yellow

Procedure: Procedure: 100 mg of the catalyst was charged in the microreactor to this 500 mg (3.42 * 10⁻³ mol) benzyledene acetone was added as solution in 2 ml tetrahydrofuran. Microreactor was flushed with argon and heated to 90 °C. Magnetic agitation was started. After attaining temperature reactor was pressurized with 500-psi hydrogen. The reactor was maintained under these conditions for 24 hours. Reaction was stopped by cooling reactor to 0°C and depressurizing. Reactor was opened and liquid was analyzed by gas chromatograph. Total benzyledene acetone was converted. Catalyst was recovered by washing reactor with several portions of tetrahydrofuran combined fractions were centrifuged to recover catalyst. Recovered catalyst was washed with water, methanol and diethyl ether and catalyst was dried under vacuum. This catalyst was recycled to obtain equivalent activity and selectivity. Catalyst was recycled to obtain equivalent conversion and selectivity.

Example 456

Nitrotolune hydrogenation

Catalyst specifications:

Catalytic entity	Ru(Cl)(μCl)(TPPTS) ₂ (10 ⁻⁶ mols)
Additive	TPPTS (6 * 10 ⁻⁶ mols)/ 500 mg sodium alginate
Support	Titania 5 g
Group IIA metal salt	Strontium chloride saturated solution
Method of preparation	coprecipitation
Color of catalyst	Pale yellow

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Procedure: 100 mg of the catalyst was charged in the microreactor to this 500 mg (3.649 * 10⁻³ mol) o-nitro toluene was added as solution in 2 ml ethyl acetate. Microreactor was flushed with argon and heated to 90 °C, magnetic agitation was started. After attaining temperature reactor was pressurized with 500-psi hydrogen. The reactor was maintained under these conditions for 24 hours. Reaction was stopped by cooling reactor to 0 °C and depressurizing. Reactor was opened and liquid was analyzed by gas chromatograph. Total o-Nitrotolune was converted to 0 toludene.

Catalyst was recovered by washing reactor with several portions of ethyl acetate combined fractions were centrifuged to recover catalyst. Recovered catalyst was washed with water, methanol and diethyl ether and catalyst was dried under vacuum. This catalyst was recycled to obtain equivalent activity and selectivity. Catalyst was recycled to obtain equivalent conversion and selectivity.

Example 457

Hydrogenation o-chloro nitro benzene

Catalyst specifications:

Catalytic entity	Ru(Cl)(μCl)(TPPTS) ₂ (10 ⁻⁶ mols)
Additive	TPPTS (6 * 10 ⁻⁶ mols)/ 500 mg sodium
	alginate
Support	Titania 5 g
Group IIA metal salt	Strontium chloride saturated solution
Method of preparation	Coprecipitation
Color of catalyst	Pale yellow

Procedure: 100 mg of the catalyst was charged in the microreactor to this 500 mg (3.952 * 10⁻³ mol) o-chloro nitrobenzene was added as solution in 2 ml ethyl acetate. Microreactor was flushed with argon and heated to 90 °C. Magnetic agitation was started. After attaining temperature reactor was pressurized with 500-psi hydrogen. The reactor was maintained under these conditions for 24 hours. Reaction was stopped by cooling reactor to 0 °C and depressurizing. Reactor was opened and liquid was analyzed by gas chromatograph. Total o-chloro nitrobenzene was converted to o chloro aniline.

Catalyst was recovered by washing reactor with several portions of ethyl acetate combined fractions were centrifuged to recover catalyst. Recovered catalyst was washed with water, methanol and diethyl ether and catalyst was dried under vacuum. This catalyst was recycled to obtain equivalent activity and selectivity. Catalyst was recycled to obtain equivalent conversion and selectivity.

Example 458

Iodobenzene and methyl acrylate

Catalyst specifications:

Catalytic entity	Pd ⁰ (P(Ph _m SO ₃) ₃) ₃ (10 ⁻⁵ mols)
Additive	P(Ph _m SO ₃) ₃ (4 * 10 ⁻⁵ mols)
Support	Charcoal 1 g
Group IIA metal salt	Barium nitrate saturated solution
Method of preparation	Deposition precipitation
Color of catalyst	Black

Procedure: procedure: procedure: procedure: 100 mg of the catalyst was charged in the microreactor to this 1 mg tetrabutyl ammonium hydroxide, 0.41 mg (5 * 10⁻³) sodium acetate 0.5 g (5 * 10⁻³) ethyl acrylate and 0.509 g (2.5 * 10⁻³) iodobenzene was added as solution in 5 ml toluene. Microreactor was flushed with argon and heated to 90 °C, magnetic agitation was started after attaining temperature. The reactor was maintained under these conditions for 48 hours. Reaction was stopped by cooling reactor to 0 °C and depressurizing. Reactor was opened and liquid was analyzed by gas chromatograph. 86 % ethyl acrylate was converted to products. Catalyst was recovered by washing reactor with several portions of toluene combined fractions were centrifuged to recover catalyst. Recovered catalyst was washed with water, methanol and diethyl ether and catalyst was dried under vacuum. This catalyst was recycled to obtain equivalent activity and selectivity. Catalyst was recycled to obtain equivalent conversion and selectivity.

Observation color of the recovered catalyst was dark yellow.

Example 459

Iodobenzene and acrylonitrile

Catalyst specifications:

Catalytic entity	Pd ⁰ (P(Ph _m SO ₃ ⁻) ₃) ₃ (10 ⁻⁵ mols)
Additive	P(Ph _m SO ₃) ₃ (4 * 10 ⁻⁵ mols)
Support	Charcoal 1 g
Group IIA metal salt	Barium nitrate saturated solution
Method of preparation	Deposition precipitation
Color of catalyst	Black

Procedure: procedure: procedure: 100 mg of the catalyst was charged in the microreactor to this 1 mg tetrabutyl ammonium hydroxide, 0.41 g sodium acetate, 0.265 g $(5 * 10^{-3})$ acrylonitrile and 0.509 g $(2.5 * 10^{-3})$ iodobenzene was added as a solution in 5

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ml toluene. Microreactor was flushed with argon and heated to 90 °C. Magnetic agitation was started after attaining temperature. The reactor was maintained under these conditions for 48 hours. Reaction was stopped by cooling reactor to 0 °C and depressurizing. Reactor was opened and liquid was analyzed by gas chromatograph. 90 % acrylonitrile was converted to products.

Catalyst was recovered by washing reactor with several portions of toluene combined fractions were centrifuged to recover catalyst. Recovered catalyst was washed with water, methanol and diethyl ether and catalyst was dried under vacuum. This catalyst was recycled to obtain equivalent activity and selectivity. Catalyst was recycled to obtain equivalent conversion and selectivity.

Observation color of the recovered catalyst was dark yellow.

Example 460

Iodobenzene and styrene

Catalyst specifications:

Catalytic entity	Pd ⁰ (P(Ph _m SO ₃ ⁻) ₃) ₃ (10 ⁻⁵ mols)
Additive	P(Ph _m SO ₃) ₃ (4 * 10 ⁻⁵ mols)
Support	Charcoal 1 g
Group IIA metal salt	Barium nitrate saturated solution
Method of preparation	Deposition precipitation
Color of catalyst	Black

Procedure: 1g of the catalyst was charged in the microreactor to this 5 mg tetrabutyl ammonium hydroxide, 0.66 mg potassium carbonate 0.5g (4.8 * 10⁻³ mol) styrene and 1.957 g (9.6 * 10⁻³ mol) iodobenzene was added as solution in 10 ml toluene. Microreactor was flushed with argon and heated to 90 °C. Magnetic agitation was started after attaining temperature. The reactor was maintained under these conditions for 76 hours. Reaction was stopped by cooling reactor to 0 °C. Reactor was opened and liquid was analyzed by gas chromatograph. 44 % styrene was converted to stilbene.

Catalyst was recovered by washing reactor with several portions of toluene combined fractions were centrifuged to recover catalyst. Recovered catalyst was washed with water, methanol and diethyl ether and catalyst was dried under vacuum. This catalyst was recycled to obtain equivalent activity and selectivity. Catalyst was recycled to obtain equivalent conversion and selectivity.

Example 461

Iodobenzene and ethylene

Catalyst specifications:

Catalytic entity	Pd (Ac) ₂ (P(o Me-Ph _m SO ₃) ₃) ₂ (10 ⁻⁵ mols)
Additive	(P(o Me-Ph _m SO ₃) ₃) ₂ (4 * 10 ⁻⁵ mols)
Support	Charcoal 1 g
Group IIA metal salt	Barium nitrate saturated solution
Method of preparation	Deposition precipitation
Color of catalyst	Black

Procedure: 100 mg of the catalyst was charged in the microreactor to this 1 mg tetrabutyl ammonium hydroxide, 0.41 g sodium acetate and 0.509 g (2.5 * 10⁻³) iodobenzene was added as solution in 10 ml acetonitrile. Microreactor was flushed with nitrogen and heated to 120°C. Reactor was pressurized with ethylene and magnetic agitation was started. The reactor was maintained under these conditions for 76 hours. Reaction was stopped by cooling reactor to 0°C. Reactor was depressurized by venting gas in the reactor. Reactor was opened and liquid was analyzed by gas chromatograph. 30 % iodobenzene was converted to styrene.

Catalyst was recovered by washing reactor with several portions of toluene combined fractions were centrifuged to recover catalyst. Recovered catalyst was washed with water, methanol and diethyl ether and catalyst was dried under vacuum. This catalyst was recycled to obtain equivalent activity and selectivity. Catalyst was recycled to obtain equivalent conversion and selectivity.

Example 462

Bromo benzene and o tolyl boronic acid

20 Catalyst specifications:

Catalytic entity	Pd ⁰ (P(Ph _m SO ₃ ') ₃) ₃ (10 ⁻⁵ mols)
Additive	P(Ph _m SO ₃ ⁻) ₃ (4 * 10 ⁻⁵ mols)
Support	Charcoal 1 g
Group IIA metal salt	Barium nitrate saturated solution
Method of preparation	Deposition precipitation
Color of catalyst	Black

Procedure: Thoroughly dried 500-ml flask was equipped with thermometer, magnetic

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stirrer bar, condenser, addition funnel and two-way valve. To the flask was placed 12.2 g (0.5 atom) of magnesium turnings. Assembly was thoroughly evacuated through two-way valve and nitrogen was filled and magnesium was stirred for 6 hours. To this was added a crystal of iodine, 100 ml of tetrahydrofuran distilled over sodium benzophenone ketyl and 200 µl of 1,2 dibromoethane. After surface of magnesium turns shiny white 78.5-g (0.5 mol) bromobenzene was added with such a rate that temperature raises to boiling. Reaction was continued with intermittent cooling of the flask by removing heater-stirrer. Once reaction subsides mixture was refluxed until magnesium was dissolved. Reaction mixture was transferred to Shlenk tube plugged with glass wool.

m-tolyl boronic acid

To a thoroughly dried flanged flask attached with sealed mechanical stuffing box and a dropping funnel and reflux condenser attached with fused calcium chloride guard tubes was assembled. Temperature of the flask was brought to -75° C with acetone and liquid nitrogen. To the flask was added 40.5 g of tributyl borate in 150-ml ether. With fairly rapid stirring add solution of o tolyl magnesium bromide without letting temperature to rise above -70° C continue stirring for three hours at same temperature. Temperature of the flask was maintained to 5 $^{\circ}$ C with ice bath for 12 hours. This reaction mixture was added to chilled 10 % sulfuric acid 150 ml. Extract with ether and evaporated to this was added 100 ml water and basified with NaOH to slightly alkaline. Acidify and extract with boiling water and collected as crystalline material (5 g).

1g of the catalyst was charged in the round-bottomed flask attached to reflux condenser, magnetic bar was added to round bottomed flask. To this 1-mg tetrabutyl ammonium hydroxide, 75 mg. sodium carbonate, 0.136-g (10⁻³ mols) o- tolyl boronic acid and 0.224 g (1.1 * 10⁻³) iodobenzene was added as solution in 20ml toluene. Assembly was flushed with nitrogen and heated to 90 °C. The reactor was maintained under these conditions for 24 hours. Reaction was stopped by cooling reactor to 0 °C. Liquid was analyzed by gas chromatograph. 95 % iodobenzene was converted to 2 methyl 1, 1' biphenyl.

Catalyst was recovered by washing reactor with several portions of toluene combined fractions were centrifuged to recover catalyst. Recovered catalyst was washed with sodium carbonate, water, methanol and diethyl ether and catalyst was dried under vacuum. This catalyst was recycled to obtain equivalent activity and selectivity. Catalyst was recycled to obtain equivalent conversion and selectivity.

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Example 463

Phenyl magnesium bromide and iodobenzene

Catalyst specifications:

Catalytic entity	NiCl ₂ .dppe ts (5 * 10 ⁻³ atom of nickel)
Additive	Dppe 0.806 mg
Support	Silica davisil
Group IIA metal salt	Barium nitrate saturated solution
Method of preparation	Fluidized bed
Color of catalyst	Pale blue
Moisture content	Not detectable by Karl Fischer

Procedure:

Preparation of Grignard reagent

Thoroughly dried 500-ml flask was equipped with thermometer, magnetic stirrer bar, condenser, addition funnel and two-way valve. To the flask was placed 0.61 g (0.025 atom) g of magnesium turnings. Assembly was thoroughly evacuated through two-way valve and nitrogen was filled and magnesium was stirred for 6 hours. To this was added a crystal of iodine, 50 ml of tetrahydrofuran distilled over sodium benzophenone ketyl and 200 µl of 1,2 dibromoethane. After surface of magnesium turns shiny white 3.922-g (0.025 mol) bromobenzene in 20-ml thf was added with such a rate that temperature raises to boiling. Reaction was continued with intermittent cooling of the flask by removing heater-stirrer. Once reaction subsides mixture was refluxed until magnesium was dissolved. Reaction mixture was transferred to Shlenk tube plugged with glass wool.

Thoroughly dried 250 ml flask was equipped with thermometer, magnetic stirrer bar, condenser, addition funnel and two way valve was charged with 5 g of catalyst, 50 ml tetrahydrofuran freshly distilled over sodium benzophenone ketyl of blue color. Assembly was filled with nitrogen as described earlier. To this was added 5.09g (0.025 mol of iodobenzene) 60 ml Grignard reagent as prepared previously was charged in addition vessel. Grignard reagent was slowly added to contents of the flask which was previously cooled to 5°C. Temperature of the flask was maintained at 5 °C for 24 h. Reaction mixture was cooled to room temperature to which was slowly added 20 ml water followed by saturated 20 ml ammonium chloride. Resulting suspension was fluttered to remove solids and subsequently washed thoroughly with tetrahydrofuran and water. Filtrates were extracted with dichloromethane to obtain biphenyl in 89 % yield residue left after filtration was washed with water, tetrahydrofuran and ether and dried under vacuum.

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Example 464

Isobutyl magnesium bromide and iodobenzene

Preparation of catalysts: catalyst was previously dried by extraction with boiling THF over sodium wire followed by vaccume and stored over phosphorus pentoxide catalyst specifications:

Catalytic entity	NiCl ₂ .dppe ts (5 * 10 ⁻³ atom of nickel)
Additive	Dppe 0.806 mg
Support	Silica davisil 5 g
Group IIA metal salt	Barium nitrate saturated solution
Method of preparation	Fluidized bed
Color of catalyst	Pale blue
Moisture content	Not detectable by Karl Fischer

Procedure: preparation of Grignard reagent

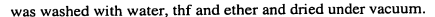
Thoroughly dried 500-ml flask was equipped with thermometer, magnetic stirrer bar, condenser, addition funnel and two-way valve. To the flask was placed 2.44 g (0.1 atom) of magnesium turnings. Assembly was thoroughly evacuated through two-way valve and nitrogen was filled and magnesium was stirred for 6 hours. To this was added a crystal of iodine, 50 ml of tetrahydrofuran distilled over sodium benzophenone ketyl and 200 µl of 1,2 dibromoethane. After surface of magnesium turns shiny white 9.2-g (0.1 mol) isobutyl bromide was added with such a rate that temperature raises to boiling. Reaction was continued with intermittent cooling of the flask by removing heater-stirrer. Once reaction subsides mixture was refluxed until magnesium was dissolved. Reaction mixture was transferred to Shlenk tube using canula plugged with glass wool.

Thoroughly dried 500 ml flask was equipped with thermometer, magnetic stirrer bar, condenser, addition funnel and two way valve was charged with 5 g of catalyst, 50-ml tetrahydrofuran freshly distilled over sodium benzophenone ketyl of blue color. Assembly was filled with nitrogen as described earlier. To this was added 20.39 g (0.1 mol of iodobenzene) 50 ml Grignard reagent as prepared previously was charged in addition vessel. Grignard reagent was slowly added to contents of the flask which were cooled to 0°C. Temperature of the flask was raised to 50-0°C temperatures. Reaction mixture was cooled to room temperature to, which was slowly added 25-ml water followed by 25 ml saturated ammonium chloride. Resulting suspension was filtered to remove solids and subsequently washed thoroughly with tetrahydrofuran and water. Filtrates were extracted with dichloromethane to obtain isobutyl benzene in 92 % yield residue left after filtration

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Example 465

Allylation of aryl boronates

Catalyst specifications:

Catalytic entity	NiCl ₂ .(tppts) ₂ (96.5 mg(10 ⁻⁴ mols)
Additive	tppts(83.6 mg(2 * 10 ⁻⁴ mols)
Support	Silica davisil 2 g
Group IIA metal salt	Barium nitrate saturated solution
Method of preparation	Fluidized bed
Color of catalyst	White with blue ting
Moisture content	Not detectable by Karl Fischer

Procedure: to a 2 gm of catalyst was added 20 ml of tetrahydrofuran, to this solution was added 5 ml of cold solution containing 2.9 mmol of phenyl lithium. The mixture was cooled to 0 °C under stirring and to this 0.42 ml (3.62 mmol) B(OCH₃)₃ was slowly added followed by 1.44 mmol allyl methyl carbonate. Temperature was then raised to 60 °C and reaction was continued for 12 hours. Liquid was separated from solid catalyst and poured in to mixture of 20 ml hexane and 20 ml of saturated ammonium chloride. Organic layer indicated formation of 3-phenyl propene.

Recovered catalyst was washed with saturated bicarbonate, tetrahydrofuran and diethyl ether and recycled after drying in vaccume

Example 466

Hexene isomarization

Catalyst specifications:

Catalytic entity	Rh ⁺ (ClO ₄ -)(tppts) ₃
Additive	tppts
Support	silica
Group IIA metal salt	Saturated barium nitrate solution
Method of preparation	Deposition precipitation
Color of catalyst	Pale yellow

Procedure: Isomarization was carried out according following procedure. 5 gm of catalyst was charged in microreactor, which was subsequently flushed with nitrogen and charged with degassed mixture of 5 g 1-hexene (92 % purity), and 45-ml cyclohexane. Temperature

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of the reactor was raised to 100 °C and maintained for 76 hours. Conversion of 1 hexene was 73 %. And two isomarized products were observed. Catalyst was centrifuged and liquid was separated. Catalyst was repeatedly washed by cyclohexane. Isolated catalyst was recycled under identical conditions to provide equivalent yields.

Example 467

N, N – Diethylneryl amine isomarization

Catalyst specifications:

Catalytic entity	Rh ⁺ (ClO ₄ ⁻)(binapts) ₃ (10 ⁻⁶ mol)
Additive	Binapts (5 * 10 ⁻⁶ mol)
Support	Silica 2g
Group IIA metal salt	Saturated barium nitrate solution
Method of preparation	Deposition precipitation
Color of catalyst	Pale yellow

Procedure: catalyst drying

Isomarization was carried out as per procedure adopted from (Helvetica Chemica Acta vol.71, (1988) 897-920) modified to suit solid catalyst. 2 gm of catalyst was charged in Fischer-porter bottle and evacuated. Bottle was subsequently flushed with nitrogen and charged with degassed mixture of 11.37 g (50 mmol) (purity by area % on gc 92) N, N – Diethylneryl amine and 50 ml dry tetrahydrofuran. Temperature of the bottle was raised to 80°C and maintained for 76 hours. Catalyst was centrifuged and liquid was separated. Repeated washings of catalyst by tetrahydrofuran were combined and evaporated to obtain pale yellow oil which was dissolved in 50 % acetic acid in water 50 ml at 0°C stirred for 10 min and 50 ml hexane was added and liquid was stirred for 30 min at ambient temperature. Hexane layer was separated and aqueous layer was washed with hexane hexane extract was washed with sat bicarbonate solution. Fractionation of extract provided 7.35 g (S) citronellol 90 % based on N, N – Diethylneryl amine. Optical purity of this material found by polarimetry c = 5, CHCl₃, lamp D at 20°C was 98 %, isolated catalyst was recycled under identical conditions to provide equivalent yields.

Example 468

25 1.4-diacetoxy butene isomarization

Catalytic entity	ClO ₄ Rh	⁺ [P(Ph _m SO ₃) ₃] ₃ (10 ⁻⁶)
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Additive	P(Ph _m SO ₃ ⁻) ₃ (6 * 10 ⁻⁶)
Support	Bentonite 5 g
Group IIA metal salt	Strontium chloride
Method of preparation	Deposition precipitation in coating pan
Color of catalyst	Pale yellow
Metal content	

Procedure: isomarization was carried out according following procedure. 100 mg of catalyst was charged in microreactor which was subsequently flushed with nitrogen and charged with degassed mixture of 50 mg (3 * 10 ⁻⁴mol) 1,4-diacetoxy butene and 2 ml toluene. Temperature of the reactor was raised to 100 ⁰C and maintained for 76 hours. Catalyst was centrifuged and liquid was separated. Repeated washings of catalyst by toluene. Conversion of 1,4-diacetoxy butene was 57 %. Isolated catalyst was recycled under identical conditions to provide equivalent yields.

Example 469

Hexene wacker

Catalyst specifications:

Palladium acetate/bypyridyldisulfonate	22.4.mg :55 mg
bypyridyldisulfonate	218.4 mg(0.4 mmol)
Support	Bentonite 2 gm
Group IIA metal salt	Strontium chloride
Method of preparation	Deposition precipitation by azeotropic removal of water
Color of catalyst	Light yellow orange
Metal content	0.46 miliatoms pre gm

Procedure: 100 mg of the catalyst was charged in the microreactor to this 1 ml hexene was added as 20 % v/v solution in hexane. Microreactor was pressurized with air (450 psi) and heated to 90°C. Magnetic agitation was started after attaining temperature. The reactor was maintained for 24 hours. Reaction was stopped by cooling reactor to 0°C and depressurizing. Reactor was opened and liquid was analyzed by gas chromatograph. 36% cyclohexene was converted to products with 90 % selectivity for hexane 2 one. Remainder products were estimated to be isomarized olefins and some unidentified products.

Catalyst was recovered by washing reactor with several portions of cyclohexane combined fractions were centrifuged to recover catalyst. Recovered catalyst was fortified with water

to contain 50 % water by weight. This catalyst was recycled to obtain equivalent activity and selectivity.

Observation color of the recovered catalyst was dark yellow.

Example 470

5 Decene wacker

Catalyst specifications:

Palladium acetate/bypyridyldisulfonate	22.4.mg :55 mg
bypyridyldisulfonate	218.4 mg(0.4 mmol)
Support	Bentonite 2 gm
Group IIA metal salt	Strontium chloride
Method of preparation	Deposition precipitation by azeotropic removal of water
Color of catalyst	Light yellow orange
Metal content	0.46 miliatoms pre gm

Procedure: procedure: 100 mg of the catalyst was charged in the microreactor to this 1 ml decene was added as 20 % solution in hexene. Microreactor was pressurized with air (450 psi) and heated to 90 °C. Magnetic agitation was started after attaining temperature. The reactor was maintained for 24 hours. Reaction was stopped by cooling reactor to 0°C and depressurizing. Reactor was opened and liquid was analyzed by gas chromatograph. 19 % decene was converted to products with 87% selectivity for decane 2 one. Remainder products were estimated to be isomarized olefins and some unidentified products.

Catalyst was recovered by washing reactor with several portions of cyclohexane combined fractions were centrifuged to recover catalyst. Recovered catalyst was fortified with water to contain 50 % water by weight. This catalyst was recycled to obtain equivalent activity and selectivity.

Observation color of the recovered catalyst was dark yellow.

Example 471

20 Cyclohexene wacker

Palladium acetate/bypyridyldisulfonate	22.4.mg :55 mg
bypyridyldisulfonate	218.4 mg(0.4 mmol)
Support	Bentonite 2 gm
Group IIA metal salt	Strontium chloride

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Method of preparation	Deposition precipitation by azeotropic
	removal of water
Color of catalyst	Light yellow orange
Metal content	0.46 miliatoms per gm

Procedure: 100 mg of the catalyst was charged in the microreactor to this 1 ml cyclohexene was added as 20 % solution in hexene. Microreactor was pressurized with air (450 psi) and heated to 90 °C. Magnetic agitation was started after attaining temperature. The reactor was maintained for 24 hours. Reaction was stopped by cooling reactor to 0 °C and depressurizing. Reactor was opened and liquid was analyzed by gas chromatograph. 7% cyclohexene was converted to products with 30 % selectivity for cyclohexanone. Remainder products were not estimated.

Catalyst was recovered by washing reactor with several portions of cyclohexane combined fractions were centrifuged to recover catalyst. Recovered catalyst was fortified with water to contain 50 % water by weight. This catalyst was recycled to obtain equivalent activity and selectivity.

Observation color of the recovered catalyst was dark yellow.

Example 472

Styrene epoxidation

Preparation of catalysts: Ph(CH₂)-N⁺(Ph_mSO₃⁻)₃ .OH⁻ was heterogenized as described according method described earlier as deposition precipitation in coating pan. Solid was suspended in water to which three mole equivalent of OH present (estimated by titration with standard acid) suspension was stirred for 4 hours at 70 °C. Solid was recovered and extracted with water for 12 hours followed by drying in vacuum.

Catalytic entity	Ph(CH ₂)-N ⁺ (Ph _m SO ₃ ⁻) ₃ . MO ₄ ⁻² (1 mili eq of MO_4 ⁻²)
additive	Sodium meta silicate 1 gm
Support	keisulghur 5 gm
Group IIA metal salt	Barium hydroxide saturated solution in water.
Method of preparation	Deposition precipitation in coating pan
Moisture content	20 % by weight.

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Procedure: 5 g of the catalyst was charged in the 500 ml glass reaction vessel equipped with mechanical stirrer, thermometer pocket and addition vessel, to this 30 g (0.29 mol) styrene in 75 ml acetic acid was added. Reaction vessel was cooled to 5 °C with circulating fluid cryostat. Agitation was started after attaining temperature. 30ml 34 % hydrogen peroxide was added over the period of 30 min. temperature and agitation of the reaction vessel was maintained for 24 hours. Reactor was opened and liquid was analyzed by gas chromatograph. 72% styrene was converted to products with 89 % selectivity for styrene oxide. Remainder products were not estimated.

Catalyst was recovered by washing reactor with several portions of acetic acid combined fractions were centrifuged to recover catalyst. Recovered catalyst was washed with methanol, ether and dried. This catalyst was recycled to obtain equivalent activity and selectivity.

Example 473

Chlorophenol oxidation

Preparation of catalysts: catalyst was prepared according to method described as precipitation in fluidized bed.

Catalyst specifications:

Catalytic entity	Iron(III) pthalocynine-4,4',4'',4'''-
	tetrasulfonic acid tetrasodium salt as
	compound with oxygen 30.2 mg (3 * 10 ⁻³
	mols)
Additive	Sodium silicate 500 mg
Support	Fullers earth 2g
Group IIA metal salt	Barium nitrate saturated solution in water
Method of preparation	Fluidized bed drier
Color of catalyst	Pale blue
Moisture content	10 % by weight

Procedure: to a 50 ml round bottomed flask attached with a reflux condenser was added 5 ml acetonitrile and 15 ml 0.1 M acetate buffer of pH 7 to which was added 181 mg (10⁻³ mols) of trichloro phenol. 2.5g catalyst was added and suspension was stirred with magnetic needle. Temperature of the suspension was raised to 60 °C. To this 0.3 ml 34%

 H_2O_2 in water was added to above suspension. The reaction mixture was continued for 5 hours. Total disappearance of trichlorophenol was observed and chloride ions were detected in solution with silver nitrate solution.

Example 474

5 Condensation of diethylfumarate and diethylmalonate to propane-1,1,2,3 tetracarboxylate Catalyst specifications:

Catalytic entity	Ph(CH ₂)-N ⁺ (Ph _m SO ₃ ⁻) ₃ . EtO ⁻ (1 milieq of
	EtO ⁻)
additive	Sodium meta silicate 1 gm
Support	Charcoal 5 gm
Group IIA metal salt	Barium hydroxide
Method of preparation	Deposition precipitation

Procedure: in a 100-ml flask fitted with an efficient reflux condenser, magnetic stirrer bar and dropping funnel was charged 5 gm of catalyst. With stirring 1.6 g (0.001 mol) diethylmalonate and 25 ml dry ethanol was added. Reaction mixture was warmed and 1.4 gm (0.0081 mol) diethyl fumarate was added. Mixture was refluxed for 8 hours. Reaction mixture was cooled and suspension was centrifuged to recover catalyst. Analysis of reaction liquid indicated 90 % conversion for diethylmalonate. Product was distilled under vacuum 8 mm at 180-190°C to obtain propane-1,1,2,3 tetracarboxylate 85% yield.

Catalyst was recovered and washed with ethanol, diethyl ether and dried under vacuum.

Catalyst was recycled to obtain equivalent activity.

Example 475

Diethymalonate and formaldehyde condensation to tetraethylpropane-1,1,3,3-tetracarboxylate

Catalytic entity	Ph(CH ₂)-N ⁺ (Ph _m SO ₃ ⁻) ₃ . HO ⁻ (1 milieq of
	HO ⁻)
additive	Sodium meta silicate 1 gm
Support	Charcoal 5 gm
Group IIA metal salt	Barium hydroxide
Method of preparation	Deposition precipitation
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Procedure: mixture of 1.6 g (0.001 mol) of distilled diethyl malonate, 25 ml ethanol and 0.4 g (5 * 10 ⁴ mol) of 40 % formaldehyde contained in 50 ml round bottomed flask was cooled to 0 ⁰C and 5 gm of catalyst was added and mixture was stirred at room temperature for 24 hours and then refluxed for 12 hours. Suspension was centrifuged to recover catalyst. Analysis of reaction liquid indicated 80 % conversion of diethyl malonate. Liquid was evaporated and extracted with diethyl ether. Extract was dried with sodium sulfate. Catalyst was washed with ethanol, diethyl ether and dried under vaccume. Catalyst was recycled to obtain equivalent activity.

Example 476

Condensation of acetone and chloroform to chlorbutol Catalyst specifications:

Catalytic entity	Ph(CH ₂)-N ⁺ (Ph _m SO ₃) ₃ . OH (1 milieq of
	OH) 0.659 mg
additive	Sodium meta silicate 1 gm
Support	Charcoal 5 gm
Group IIA metal salt	Barium hydroxide
Method of preparation	Deposition precipitation

Procedure: 100 ml round bottomed flask equipped with reflux condenser was charged with 5 gm of catalyst to which solution of 1.19 g (0.01 mol) chloroform was charged as solution in 25 ml solution in acetone. Magnetic stirrer bar was added in the reaction mixture reaction mixture was stirred at ambient temperature for 24 hours. Suspension was centrifuged to recover catalyst. Analysis of reaction liquid indicated total conversion of chloroform. Catalyst was washed with ethanol, diethyl ether and dried under vacuum. Catalyst was recycled to obtain equivalent activity.

Example 477

20 Condensation of benzaldehyde and acetonitrile to cinnamonitrile.

Catalytic entity	Ph(CH ₂)-N ⁺ (CH ₂ Ph _m SO ₃ ⁻) ₃ . OH ⁻ (1 milieq
	of OH ⁻) 0.701 mg
Additive	Sodium meta silicate 1 gm
Support	Charcoal 5 gm
Group IIA metal salt	Barium hydroxide
Method of preparation	Deposition precipitation

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Procedure: 250 ml round bottomed flask equipped with reflux condenser was charged with 5 gm of catalyst to which solution of 10.6 g (0.1 mol) benzaldehyde was charged as solution in 100 ml solution in acetonitrile. Magnetic stirrer bar was added in the reaction mixture reaction mixture was stirred at reflux temperature for 24 hours. Suspension was centrifuged to recover catalyst. Analysis of reaction liquid indicated 88 % conversion of benzaldehyde and 98 % selectivity for cinnamonitrile. Cinnamonitrile was recovered by fractional distillation 80 % based on benzaldehyde

Catalyst was washed with ethanol, diethyl ether and dried under vacuum. Catalyst was recycled to obtain equivalent activity.

Example 478

Condensation of benzaldehyde and acetone to benzayledene acetone Catalyst specifications:

Catalytic entity	Ph(CH ₂)-N ⁺ (CH ₂ Ph _m SO ₃ ⁻) ₃ . OH ⁻ (1 milieq
·	of OH') 0.701 mg
additive	Sodium meta silicate 1 gm
Support	Charcoal 5 gm
Group IIA metal salt	Barium hydroxide
Method of preparation	Deposition precipitation

Procedure: 250 ml round bottomed flask equipped with reflux condenser was charged with 5 gm of catalyst to which solution of 10.6 g (0.1 mmol) benzaldehyde was charged as solution in 100 ml solution in acetone. Magnetic stirrer bar was added in the reaction mixture reaction mixture was stirred at ambient temperature for 24 hours. Suspension was centrifuged to recover catalyst. Analysis of reaction liquid indicated 77 % conversion benzaldehyde. Catalyst was washed with ethanol, diethyl ether and dried under vaccume. Catalyst was recycled to obtain equivalent activity.

Example 479

Condensation of butaraldehyde to 2ethyl hexenal

Catalyst specifications:

Catalytic entity	Ph(CH ₂)-N ⁺ (Ph _m SO ₃) ₃ . OH (1 milieq of OH)
	0.659 mg
Additive	Sodium meta silicate 1 gm
Support	Charcoal 5 gm
Group IIA metal salt	Barium hydroxide
Method of preparation	Deposition precipitation

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Procedure: 100 ml round bottomed flask equipped with reflux condenser was charged with 5 gm of catalyst to which solution of 7.2 g (0.1 mmol) butaraldehyde was charged as solution in 50 ml solution in toluene. Magnetic stirrer bar was added in the reaction mixture reaction mixture was refluxed for 24 hours. Suspension was centrifuged to recover catalyst. Analysis of reaction liquid indicated 90 % conversion of butaraldehyde. Catalyst was washed with ethanol, diethyl ether and dried under vacuum. Catalyst was recycled to obtain equivalent activity.

Example 480 Iodobenzene phosphination catalyst specifications:

Catalytic entity	NiCl ₂ .(bisdiphenylphosphinoethane
	tetrasulfonate) [1 miliatom of nickel]
Additive	bisdiphenylphosphinoethane tetrasulfonate
	1 gm.
support	γ- alumina 5 gm
Cured with	Barium nitrate
Method of preparation	Deposition precipitation in coating pan
Color of catalyst	Pale yellow
Metal content of prepared catalyst	0.93 miliatom of nickel

Procedure: round bottomed flask equipped with reflux condenser was charged with catalyst 5 gm. To this solution of diphenyl phosphine 1 ml (5.75 mmol) in 30 ml dry degassed dimethylformamide was added at room temperature. Suspension was degassed with repetitive vaccume and argon flushing. After heating to 100 °C for 30 min. 10 mmol (2.04 gm.) iodobenzene and 20 mmol (2.25 gm) diazabicyclooctane in 30 ml dimethylformamide was added and resulting solution was maintained at 100 °C. Three additional portions of 1-ml diphenyl phosphine each were added at 12-hour interval thereafter. Reaction was continued for 76 hours. Reaction was stopped by cooling flask to room temperature. Catalyst was recovered by centrifugation and washed by dimethylformamide. Filtrates were combined and evaporated to obtain sticky residue, which was diluted with 50-ml tetrahydrofuran. Solution was analyzed with 31 P NMR. Following compounds were detected triphenylphosphine, triphenylphosphine oxide, diphenylphosphine and diphenylphosphineoxide.

Yield of triphenylphosphine based on iodobenzene was 92%; conversion of iodobenzene was complete as confirmed by gas chromatography.

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Observation color of recovered catalyst is darker than fresh catalyst.

Catalyst recovery and recycle: The recovered catalyst was recycled under identical conditions to obtain 88% triphenylphosphine based on iodobenzene. For third recycle catalyst was refluxed with 250 mg NiCl₂.6H₂O dissolved in 50% ethanol in water for 6 hours. Catalyst was extracted with water, ethanol and dried. This catalyst was recycled to obtain 90% triphenylphosphine

Fresh catalyst under identical conditions in cyclohexane as solvent does not indicate leaching of nickel but yield of triphenyl phosphine was about 10% after 76 hours.

Example 481

Bromoanisol phosphination

Catalyst preparation: catalyst was prepared according to method described as deposition precipitation in coating pan

Catalyst specifications:

Catalytic entity	PdAc ₂ .(bisdiphenylphosphinoethane
	tetrasulfonate) [1 miliatom of palladium]
Additive	bisdiphenylphosphinoethane tetrasulfonate
	1 gm.
support	Gamma alumina 5 gm
Cured with	Barium nitrate
Method of preparation	Deposition precipitation in coating pan
Color of catalyst	Pale yellow
Metal content of prepared catalyst	0.93 miliatom of palladium

Procedure: round bottomed flask equipped with reflux condenser was charged with catalyst 5 gm. To this solution of diphenyl phosphine 1 ml (5.75 mmol) in 30-ml dry degassed dimethylformamide was added at room temperature. Suspension was degassed with repetitive vaccume and argon flushing. After heating to 100 °C for 30 min. 10 mmol (1.87 gm.) 2- bromoanisol and 20 mmol (2.25 gm) diazabicyclooctane in 30 ml dimethylformamide was added and resulting solution was maintained at 100 °C. Three additional portions of 1-ml diphenyl phosphine each were added at 12-hour interval thereafter. Reaction was continued for 76 hours. Reaction was stopped by cooling flask to room temperature. Catalyst was recovered by centrifugation and washed by dimethylformamide. Filtrates were combined and evaporated to obtain sticky residue,

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which was diluted with 50-ml tetrahydrofuran. Solution was analyzed with ³¹P NMR as described in previous example. 83 % conversion of 2 bromoanisol was observed. Quantitative estimation of phosphines was not determined.

Example 482

Duteriation of C₆H₆ to C₆ D₆
 Catalyst specifications

Catalytic entity	Diphenyl phosphinoethane tetrasulphonate
	/ RuCl ₂ COD 15 mg
Additive	Diphenyl phosphinoethane tetrasulphonate
	25 mg / sodium phosphate50 mg
Support	Bentonite 500 mg
Cured with	Saturated strontium chloride solution in
	water
Method of preparation	Fluidized bed
Color of catalyst	Pale yellow

Catalyst pretreatment: catalyst was refluxed twice with 3-ml deuterium oxide recovered with centrifugation and dried under vaccume. This was essential to remove protons on the solid support.

Procedure: 100 mg of the catalyst was charged in the microreactor equipped with external magnetic stirrer 0.01mol (0.78 g) benzene in 2 ml and deuterium oxide was added. Reaction vessel was heated to 110°C with external heating. Agitation was started after attaining temperature. Reactor was maintained at these conditions for 24 hours. Reactor was cooled to -5°C liquids were recovered and organic liquid was analyzed by NMR 88 % labeling of deuterium using chloroform as internal standard.

Catalyst was recovered by centrifugation and recovered catalyst was dried under vacuum. This catalyst was recycled to obtain equivalent activity and selectivity.

While invention has been particularly shown and described with respect to preferred embodiments thereof, it will be understood by those skilled in the art that the forgoing and the other changes in form and detail may be made without departing from the spirit and scope of the invention.